Soot formation in laminar counterflow flames

Yu Wang\textsuperscript{1,*}, Suk Ho Chung\textsuperscript{2,*}

\textsuperscript{1}School of Automotive Engineering, Wuhan University of Technology, Wuhan, P.R. China
\textsuperscript{2}Clean Combustion Research Center (CCRC), King Abdullah University of Science and Technology (KAUST), Thuwal, Saudi Arabia

* Corresponding authors

Yu Wang
School of Automotive Engineering
Wuhan University of Technology
Wuhan, P.R. China
Email: yu.wang@whut.edu.cn

Suk Ho Chung
Clean Combustion Research Center
King Abdullah University of Science and Technology
Thuwal, Saudi Arabia
Email: sukho.chung5@gmail.com

Review article
Prepared for submission to Progress in Energy and Combustion Science
Soot formation in laminar counterflow flames

Yu Wang¹,*, Suk Ho Chung²,*

¹School of Automotive Engineering, Wuhan University of Technology, Wuhan, P.R. China
²Clean Combustion Research Center (CCRC), King Abdullah University of Science and Technology (KAUST), Thuwal, Saudi Arabia

Abstract

Many practical soot-emitting combustion systems such as diesel and jet engines rely on diffusion flames for efficient and reliable operation. Efforts to mitigate soot emissions from these systems are dependent on fundamental understanding of the physicochemical pathways leading from fuel to soot in laminar diffusion flames. Existing diffusion flame–based soot studies focused primarily on over-ventilated coflow flame where the fuel gas (or vapor) issues from a cylindrical tube into a co-flowing oxidizer, and counterflow flame, where a reacting zone is established between two opposing streams of fuel and oxidizer. As a canonical diffusion flame configuration, laminar counterflow diffusion flames have been widely used as a highly controllable environment for soot research, enabling significant progress in the understanding of soot formation for several decades. In view of the possibility of fuel/oxidizer premixing in practical systems, counterflow partially premixed flames have also been studied. In the present work we intend to provide a comprehensive review of the researches on various aspects of soot formation utilizing counterflow flames. Major processes of soot formation (formation of gas phase soot precursors, soot inception and surface reactions, as well as particle-particle interactions) are examined first, with focus on the most recent (post-2010) research progress. Experimental techniques and associated challenges for the measurement of soot-related properties (some knowledge of which is helpful for full appreciation of the experimental data to be reviewed) are then introduced. This is followed by a detailed description of soot evolution in counterflow flames, which is complemented by a discussion on the similarity and differences of the sooting structures between counterflow and coflow diffusion flames. Parametric studies of the effects of fuel molecular structure, fuel additive, partial-premixing, pressure, temperature, stoichiometric mixture fraction, and residence time on soot formation in counterflow flames will then be addressed in detail. This review concludes with a summary of the knowledge and challenges gathered and demonstrated through decades of research, and an outlook on opportunities for future counterflow flame–based soot research towards a more complete understanding of soot formation and the development of novel techniques for soot mitigation in practical combustion devices.
Contents

1. INTRODUCTION ........................................................................................................................................... 6
  1.1 Motivation of soot research ......................................................................................................................... 6
  1.2 Laboratory-scale experimental configurations .......................................................................................... 9
  1.3 Structure of this review .............................................................................................................................. 14

2. PATHWAYS OF SOOT FORMATION ............................................................................................................ 15
  2.1 Gas phase precursor formation ................................................................................................................... 16
    2.1.1 Formation of first aromatic ring ............................................................................................................. 17
    2.1.2 Molecular growth beyond the first aromatic ring ................................................................................. 24
  2.2 Soot nucleation ........................................................................................................................................... 33
  2.3 Soot surface growth, particle coalescence, and agglomeration ............................................................... 42
  2.4 Soot oxidation and fragmentation ............................................................................................................. 49
  2.5 Models of soot formation .......................................................................................................................... 53

3. EXPERIMENTAL TECHNIQUES FOR SOOT RELATED MEASUREMENTS .............................................. 57
  3.1 Measurements of gas-phase species ........................................................................................................ 58
    3.1.1 Gas chromatography ............................................................................................................................ 58
    3.1.2 Mass spectrometry ............................................................................................................................... 60
    3.1.3 Optically-based in-situ methods ........................................................................................................... 64
  3.2 Soot measurements ................................................................................................................................... 70
    3.2.1 Light extinction ..................................................................................................................................... 70
    3.2.2 Spectral soot emission ........................................................................................................................... 73
    3.2.3 Laser-induced incandescence .............................................................................................................. 76
    3.2.4 Thermophoretic sampling ................................................................................................................... 79
    3.2.5 Differential mobility analysis ................................................................................................................ 81

4. SOOTING STRUCTURES AND SOOT EVOLUTION IN COUNTERFLOW FLAMES .................................... 84
  4.1 Sooting structures of CDFs and comparison with coflow diffusion flames ............................................. 84
  4.2 Influences of stoichiometric mixture fraction on sooting structures in CDFs ........................................ 92
  4.3 Influences of strain rate on sooting structures in CDFs ......................................................................... 97
  4.4 Sooting structures of partially-premixed CDFs ....................................................................................... 101

5. SOOT FORMATION IN COUNTERFLOW DIFFUSION FLAMES: PARAMETRIC STUDIES .................. 103
  5.1 Effects of fuel types on soot formation ...................................................................................................... 115
    5.1.1 Counterflow diffusion flames with single component (neat) fuels ..................................................... 116
    5.1.2 Counterflow diffusion flames with fuel mixtures ............................................................................... 123
      a) Effects of CO/H₂ addition .................................................................................................................. 123
      b) Synergistic effects of ethylene-based binary fuel mixtures ................................................................. 126
      c) Surrogates for practical fuels ........................................................................................................... 133
    5.1.3 Quantitative indices of sooting tendencies ......................................................................................... 136
  5.2 Effects of dilution and non-hydrocarbon additives on soot formation .................................................. 143
    5.2.1 Nitrogen and noble gas addition ......................................................................................................... 143
    5.2.2 Carbon dioxide and water vapor addition .......................................................................................... 152
    5.2.3 Partial-premixing ............................................................................................................................... 157
    5.2.4 Other dopants .................................................................................................................................... 165
      a) Addition of chlorinated species .......................................................................................................... 165
      b) Addition of metal-containing compounds ....................................................................................... 166
    5.3 Effects of pressure on soot formation ..................................................................................................... 170
  5.4 Effects of temperature on soot formation ................................................................................................. 180
  5.5 Effects of strain rate on soot formation ................................................................................................... 185
    5.5.1 Strain rate effects on sooting behaviors in steady CDFs .................................................................. 186
    5.5.2 Soot formation in CDFs subject to unsteady strain rates ................................................................. 190
  5.6 Various other effects .................................................................................................................................. 197
    5.6.1 Soot formation in counterflow flames of fuel sprays ....................................................................... 197
5.6.2 Effects of electric fields on soot formation in CDFs ................................................................. 200
5.6.3 Effects of soot radiation on flame characteristics ................................................................. 201
5.6.4 Interactions between soot and nitric oxides ......................................................................... 203

6. SUMMARY, CHALLENGES, AND PROSPECTS ..................................................................... 205

ACKNOWLEDGEMENTS ............................................................................................................. 210
**Nomenclature**

**Abbreviations**

<table>
<thead>
<tr>
<th>Abbreviation</th>
<th>Definition</th>
</tr>
</thead>
<tbody>
<tr>
<td>AC</td>
<td>alternative current</td>
</tr>
<tr>
<td>AFM</td>
<td>atomic force microscope</td>
</tr>
<tr>
<td>ARS</td>
<td>aromatic site</td>
</tr>
<tr>
<td>CARS</td>
<td>coherent anti-stokes Raman spectroscopy</td>
</tr>
<tr>
<td>CDF</td>
<td>counterflow diffusion flame</td>
</tr>
<tr>
<td>CI</td>
<td>compression ignition</td>
</tr>
<tr>
<td>CHC</td>
<td>chlorinated hydrocarbons</td>
</tr>
<tr>
<td>CPC</td>
<td>condensation particle counter</td>
</tr>
<tr>
<td>CRDS</td>
<td>cavity ring-down spectroscopy</td>
</tr>
<tr>
<td>DFT</td>
<td>density functional theory</td>
</tr>
<tr>
<td>DMA</td>
<td>differential mobility analysis</td>
</tr>
<tr>
<td>EI</td>
<td>electron ionization</td>
</tr>
<tr>
<td>EXP</td>
<td>experiment</td>
</tr>
<tr>
<td>FBC</td>
<td>fuel-borne catalyst</td>
</tr>
<tr>
<td>FFT</td>
<td>fast Fourier transform</td>
</tr>
<tr>
<td>FID</td>
<td>flame-ionization detector</td>
</tr>
<tr>
<td>FTIR</td>
<td>Fourier transform infra-red spectroscopy</td>
</tr>
<tr>
<td>GC</td>
<td>gas chromatography</td>
</tr>
<tr>
<td>GDI</td>
<td>gasoline direction injection</td>
</tr>
<tr>
<td>HAB</td>
<td>height above burner</td>
</tr>
<tr>
<td>HACA</td>
<td>hydrogen-abstraction-acetylene-addition</td>
</tr>
<tr>
<td>HIM</td>
<td>helium-ion microscopy</td>
</tr>
<tr>
<td>IC</td>
<td>internal combustion</td>
</tr>
<tr>
<td>ICCD</td>
<td>intensified charge coupled device</td>
</tr>
<tr>
<td>IR</td>
<td>infrared</td>
</tr>
<tr>
<td>JSR</td>
<td>jet-stirred reactor</td>
</tr>
<tr>
<td>KMC</td>
<td>kinetic Monte Carlo</td>
</tr>
<tr>
<td>LAS</td>
<td>laser absorption spectroscopy</td>
</tr>
<tr>
<td>LDV</td>
<td>laser doppler velocimetry</td>
</tr>
<tr>
<td>LE</td>
<td>light extinction</td>
</tr>
<tr>
<td>LE/LS</td>
<td>light extinction and scattering</td>
</tr>
<tr>
<td>LIE</td>
<td>laser-induced emissions</td>
</tr>
<tr>
<td>LIF</td>
<td>laser-induced fluorescence</td>
</tr>
<tr>
<td>LII</td>
<td>laser-induced incandescence</td>
</tr>
<tr>
<td>MAC</td>
<td>methyl addition/cyclization</td>
</tr>
<tr>
<td>MB</td>
<td>molecular beam</td>
</tr>
<tr>
<td>MBMS</td>
<td>molecular beam mass spectrometry</td>
</tr>
<tr>
<td>MPI</td>
<td>micropyrolysis index</td>
</tr>
<tr>
<td>MS</td>
<td>mass spectrometry</td>
</tr>
<tr>
<td>NOC</td>
<td>nanoscale organic carbon particle</td>
</tr>
<tr>
<td>NOx</td>
<td>nitic oxides</td>
</tr>
<tr>
<td>NSP</td>
<td>normalized smoke point</td>
</tr>
<tr>
<td>OESI</td>
<td>oxygen extended sooting index</td>
</tr>
<tr>
<td>PAC</td>
<td>phenyl addition/cyclization</td>
</tr>
<tr>
<td>PAH</td>
<td>polycyclic aromatic hydrocarbon</td>
</tr>
<tr>
<td>PIE</td>
<td>photo-ionization efficiency</td>
</tr>
<tr>
<td>PM</td>
<td>particulate matter</td>
</tr>
<tr>
<td>PMT</td>
<td>photomultiplier tube</td>
</tr>
<tr>
<td>PPF</td>
<td>partially-premixed flame</td>
</tr>
<tr>
<td>PSD</td>
<td>particle size distribution</td>
</tr>
<tr>
<td>REMPI</td>
<td>resonance-enhanced multi-photon ionization</td>
</tr>
<tr>
<td>RSR</td>
<td>resonantly stabilized radical</td>
</tr>
<tr>
<td>SEM</td>
<td>scanning electron microscopy</td>
</tr>
<tr>
<td>SF</td>
<td>soot formation</td>
</tr>
<tr>
<td>SFO</td>
<td>soot formation/oxidation</td>
</tr>
<tr>
<td>SMD</td>
<td>Sauter mean diameter</td>
</tr>
<tr>
<td>SMPS</td>
<td>scanning mobility particle sizer</td>
</tr>
<tr>
<td>SP</td>
<td>smoke point</td>
</tr>
<tr>
<td>SSE</td>
<td>spectral soot emission</td>
</tr>
<tr>
<td>SSI</td>
<td>sooting sensitivity index</td>
</tr>
<tr>
<td>STI</td>
<td>sooting temperature index</td>
</tr>
<tr>
<td>SVF</td>
<td>soot volume fraction</td>
</tr>
<tr>
<td>TCD</td>
<td>thermal conductivity detector</td>
</tr>
<tr>
<td>TEM</td>
<td>transmission electron microscopy</td>
</tr>
<tr>
<td>TGA</td>
<td>thermal-gravimetric analysis</td>
</tr>
<tr>
<td>TOF</td>
<td>time-of-flight</td>
</tr>
<tr>
<td>TR</td>
<td>time-resolved</td>
</tr>
<tr>
<td>TSI</td>
<td>threshold soot index</td>
</tr>
<tr>
<td>UHC</td>
<td>unburned hydrocarbon</td>
</tr>
<tr>
<td>VOA</td>
<td>vibratory orifice atomizer</td>
</tr>
<tr>
<td>UV</td>
<td>ultra-violet</td>
</tr>
<tr>
<td>VUV</td>
<td>vacuum ultra-violet</td>
</tr>
<tr>
<td>TDLAS</td>
<td>tunable diode laser absorption spectroscopy</td>
</tr>
<tr>
<td>TSI</td>
<td>threshold soot index</td>
</tr>
<tr>
<td>XRD</td>
<td>X-ray diffraction</td>
</tr>
<tr>
<td>YSI</td>
<td>yield soot index</td>
</tr>
<tr>
<td>RDG-FA</td>
<td>Rayleigh-Debye-Gans fractal aggregate</td>
</tr>
</tbody>
</table>
**Nomenclature - continued**

**Symbols**

<table>
<thead>
<tr>
<th>Symbol</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>$B$</td>
<td>Planck function</td>
</tr>
<tr>
<td>$c$</td>
<td>specific heat</td>
</tr>
<tr>
<td>$d/D$</td>
<td>diameter</td>
</tr>
<tr>
<td>$E(\tilde{m}_\lambda)$</td>
<td>light absorption function</td>
</tr>
<tr>
<td>$f_c$</td>
<td>soot volume fraction</td>
</tr>
<tr>
<td>$h$</td>
<td>Planck constant</td>
</tr>
<tr>
<td>$I$</td>
<td>light intensity</td>
</tr>
<tr>
<td>$k$</td>
<td>reaction rate constant</td>
</tr>
<tr>
<td>$K$</td>
<td>strain rate</td>
</tr>
<tr>
<td>$K_{\text{ext}}$</td>
<td>extinction coefficient</td>
</tr>
<tr>
<td>$L$</td>
<td>length of light path</td>
</tr>
<tr>
<td>$m$</td>
<td>mass</td>
</tr>
<tr>
<td>$\tilde{m}_\lambda$</td>
<td>complex refractive index</td>
</tr>
<tr>
<td>$M$</td>
<td>molecular weight</td>
</tr>
<tr>
<td>$n$</td>
<td>exponent</td>
</tr>
<tr>
<td>$N$</td>
<td>particle/molecular number concentration</td>
</tr>
<tr>
<td>$P$</td>
<td>pressure</td>
</tr>
<tr>
<td>$Q$</td>
<td>fluorescence or scattering intensity</td>
</tr>
<tr>
<td>$\dot{Q}$</td>
<td>heat loss rate</td>
</tr>
<tr>
<td>$R_g$</td>
<td>radius of gyration</td>
</tr>
<tr>
<td>$s$</td>
<td>transition line strength</td>
</tr>
<tr>
<td>$S$</td>
<td>soot surface area</td>
</tr>
<tr>
<td>$T$</td>
<td>temperature</td>
</tr>
<tr>
<td>$U$</td>
<td>internal energy</td>
</tr>
<tr>
<td>$X$</td>
<td>mole fraction</td>
</tr>
<tr>
<td>$y$</td>
<td>distance along line-of-sight chord</td>
</tr>
<tr>
<td>$Y$</td>
<td>mass fraction</td>
</tr>
<tr>
<td>$z$</td>
<td>distance from fuel nozzle</td>
</tr>
<tr>
<td>$Z$</td>
<td>mixture fraction</td>
</tr>
</tbody>
</table>

- $\alpha$ fraction of surface sites for surface reaction
- $\eta$ absorption coefficient / Stokes parameter
- $\kappa$ Boltzmann constant
- $\nu$ stoichiometric coefficient / kinematic viscosity
- $\rho$ density
- $\sigma_{\text{abs}}$ absorption cross section
- $\phi$ equivalence ratio
- $\chi$ scalar dissipation rate
- $\omega$ soot growth rate / oscillation frequency
1. Introduction

1.1 Motivation of soot research

Rapid population growth and progressive industrialization of developing countries assure rising worldwide energy demands in the coming decades: A 30% increase in energy consumption is projected from 2020 to 2040, even though continuous improvement in efficiency of energy use is anticipated [1]. Although alternative sources such as wind, solar, geothermal, and nuclear energies are increasingly important in the world energy mix, conventional petroleum-based fuels are still expected to play a dominant role in the foreseeable future. This is especially true in the transportation sector, where the society relies heavily on internal combustion (IC) engines fueled by natural gas, gasoline, kerosene, and diesel. Note that IC engines use 70% of the 86 million barrels of crude oil the world consumes every day [2]. Energy from hydrocarbons comes with a price. Apart from contributing to the greenhouse effects and thus to climate change, the use of hydrocarbon fuel also leads to the formation of various hazardous pollutants, including nitric oxide (NOx), carbon monoxide (CO), unburned hydrocarbons (UHCs) and soot. Here soot is defined as carbonaceous particles resulting from pyrolysis or incomplete combustion of hydrocarbon fuels. In terms of elemental composition, soot is largely carbon and may also contain small amounts of hydrogen and oxygen. Being very fine particles with diameters typically less than a micrometer, soot is known to be an important contributor to PM$_{2.5}$ (particulate matter with diameters smaller than 2.5 μm).

As far as reciprocating IC engines are concerned, NOx, CO, and UHC can be effectively removed using a three-way catalyst in spark-ignition (SI) engines operating with stoichiometric fuel-air mixtures. Soot emission is generally not an issue in these conventional port-injection SI engines. But with the advent of gasoline direct injection (GDI) technology (intended mainly to improve fuel efficiency), soot particles can be produced at locally rich regions as a result of charge inhomogeneity.
Although the total soot mass emitted from GDI engines may not be significant, its number density can still be tremendous, since GDI engines are likely to produce very fine soot particles [3, 4]. Compression ignition (CI) diesel engines are conventional producers of NOx and soot, and worse, a trade-off relationship exists between NOx and soot formation in the diesel combustion process. Many measures designed to reduce NOx emission result in inhibiting the oxidative removal of soot, signifying the importance of soot suppression during its formation stage. As environmental regulations on soot emission become increasingly stringent, the incentive for combustion scientists and engineers to develop innovative and effective methods for soot mitigation grows with it.

Concerns about soot emission are many and justified. Being the major condensed-phase by-product, soot originates from the incomplete combustion of hydrocarbon fuels. In most cases, the presence of soot indicates poor combustion conditions, where not all the fuel molecules can be fully oxidized to form CO$_2$ and H$_2$O for maximum thermal energy recovery. Soot particles contain chemical energy in the form of C-H and C-C bonds, which could be released as desirable thermal energy if soot formation could be avoided altogether, or if all the initially formed soot particles were fully oxidized in a later stage. Therefore, the emission of soot from practical combustion devices can be translated directly to combustion inefficiency.

More importantly, soot emission into the atmosphere has dire consequences to human health and the environment. After being inhaled, ultrafine soot particles with diameters less than 100 nm (typical for flame generated soot) can travel deeply into the lungs, deposit in the alveoli, and cross the cells to enter the circulatory system [5]. These ultrafine particles can then be translocated to other organs like the liver, heart, and brain [6], causing potential health issues such as cardiovascular diseases. Polycyclic aromatic hydrocarbons (PAHs), the main organic component of soot, have also been associated with carcinogenic and mutagenic effects [7]. Schwartz and coworkers [8, 9] demonstrated positive correlations between PM concentrations and mortality rates that reflect the serious health issues caused by soot emission.
Regarding environmental implications, soot is a major contributor to global warming, second only to CO₂ [10]. Such efficacy is believed to be caused by its influence on snow albedo as well as its atmospheric stability at high altitudes [11]. Similarly, soot deposited on ice glaciers can promote melting, raising the sea level with all the accompanying problems. Soot particles, with graphitic carbon as the primary constituent, are the biggest contributor to light absorption by atmospheric aerosols [12]. The reduction of visibility in densely populated urban areas due to the presence of soot could cause widespread societal concerns. Recent severe haze events in China, affecting nearly 800 million people, have been closely linked to fossil fuel use. It has been suggested that both primary particulate and secondary aerosol precursors from fossil fuel and biomass combustion can be controlled to minimize the negative impacts [13]. Because of its contribution as cloud condensation nuclei, soot in the upper atmosphere is also known to affect weather in the form of clouds and precipitation [14-16].

In most cases, soot is the unwanted by-product of hydrocarbon combustion and its formation needs to be suppressed as much as possible. However, there are situations in which soot production is intentional. In utility and industrial boilers, for example, released combustion heat needs to be effectively transferred to the feedwater. One important heat transfer mode in such large-scale systems is radiation, which can be significantly enhanced by the presence of soot particles [17]. Of course, these particles need to be eliminated before the flue gases are emitted to the environment. In addition, carbon black is an important raw material for the reinforcing filler in tires and the industry strives for maximum soot yield for a given quantity of hydrocarbon fuel.

The need to suppress soot emission, or to produce soot with special properties, necessitates active control of the soot formation processes. This in turn requires a fundamental understanding of the physicochemical pathways from fuel to soot. Soot formation is one of the most complex phenomena in combustion, involving complicated interactions between combustion chemistry, fluid mechanics, mass/heat transport, and particle dynamics, so that despite decades of active research, many gaps still
remain in our understanding of soot [18]. In the attempt to bridge these gaps, the underlying mechanisms of soot formation are the objectives of combustion scientists worldwide, evident from the large body of literature about soot published in recent years. For instance, soot, as a keyword, is present in the title or abstract of 45 articles published in *Combustion and Flame* in the year of 2015, accounting for more than ten percent of the 383 total articles. This number is even higher for 2016, with 51 out of the 374 articles investigating soot formation.

1.2 Laboratory-scale experimental configurations

A survey of the literature reveals that a significant portion of soot studies conducted to date are based on laboratory-scale laminar flames at atmospheric, or even lower pressures. This may seem inappropriate, at first glance, as combustion modes in most soot-emitting practical combustion devices are turbulent, and under high pressure. Such a mismatch is certainly not because soot formation in high-pressure turbulent flames can be well represented by those in laminar flames. One only needs to consider turbulence-chemistry interactions and the pressure-dependence of reaction rates to realize the drastic differences. However, the following facts limit our ability to investigate in detail the soot characteristics of high-pressure turbulent flames. First, the highly transient and inhomogeneous nature of turbulent flames presents a challenge for many soot-related experimental techniques in terms of providing sufficient temporal and spatial resolutions. Second, a high peak value and wide dynamic range of soot loadings prevent many high-pressure turbulent flames from being reliably diagnosed. Related issues include, for example, probe clogging for intrusive methods and signal attenuation/beam steering for optical methods [19]. Third, it is somewhat difficult to establish well-controlled high-pressure flames that are suitable for fundamental studies of soot formation and oxidation processes; several researchers have observed flame stability issues at high pressures [20, 21]. With an increase in pressure, the gas-solid conjugate heat transfer between flame gas and burner
nozzle becomes increasingly important and complicates the specification of flow boundary conditions [22].

In addition to the above technical constraints, the fact that physicochemical formation pathways of soot are still not fully understood—even in zero-dimensional systems—may provide enough rationale for the wide interests in soot studies with simple flow configurations. In this regard, various laboratory-scale setups have been employed, including constant volume combustion chambers [23-30], shock tubes [31-40], well-stirred reactors [41-45], burner-stabilized flat premixed flames [46-55], coflow diffusion flames [56-72], and counterflow diffusion flames (CDF) [73-92]. The conditions for soot formation can be very different among these experimental configurations and conclusions regarding the sooting processes obtained from one configuration may not be directly applicable to others. For example, the trend that soot loadings increase with the increase in flame temperature [93-96], as observed in diffusion flames, can be reversed in some premixed flames [97-99]. This can be explained (at least partly) by noting that unlike in a diffusion flame, a very rich fuel pyrolysis region, where soot precursors are formed without oxidation, usually does not exist in a premixed flame. In addition, certain fuel additives may play different roles, sometimes with opposite effects on soot formation in different reacting systems. The doping of small amounts of ethanol, or dimethyl ether, was seen to boost soot volume fractions in ethylene diffusion flames [100, 101]; but it inhibited soot formation in ethylene/air premixed flames [102, 103].

Constant volume/pressure combustion chambers are typically used to study liquid fuel spray combustion [23-30], a close representation of the combustion process in CI engines. Unfortunately, soot formation during the burning of fuel sprays depends not only on soot chemistry but on many other physical processes such as spray penetration, droplet size distribution, and velocity field of the entrained air. It is challenging to isolate these factors to obtain quantitative information on soot behaviors. Note that a shock tube can provide homogeneous, constant pressure/temperature conditions, ideal for studying the chemistry of fuel pyrolysis and subsequent soot formation. However,
the time scale of the sooting processes in shock tube experiments is typically 1~2 orders of magnitude shorter than that in flames [31]. For this reason, shock tubes are primarily used for investigation of early stage soot formation; while slower processes such as particle coagulation and aggregation cannot be effectively examined [31]. A jet-stirred reactor (JSR) is an alternative way to reach constant pressure and temperature conditions, and the residence time can be varied by changing the reactants’ flow rates. However, external heat sources are needed and typical temperature achievable in JSR experiments is only at the lower end of flame environments. Moreover, the reactants in shock tubes or JSR are highly diluted by inert gas (e.g., argon) in most cases, differing noticeably with practical flame conditions. Note that the effect of reactant dilution on soot formation can be significant [95, 104].

The details of soot formation in turbulent diffusion flames entail complex study, and most combustion systems rely on turbulent diffusion flames for operation. Fortunately, the concept of laminar flamelets could provide a link between turbulent and laminar diffusion flames [105, 106]. Fundamental investigations of the pathways from fuel molecules to soot particles in laminar diffusion flames can provide essential information toward clarifying the sooting process in practical flames.

Soot formation characteristics in laminar diffusion flames have been extensively studied in over-ventilated coflow diffusion flames, where the fuel gas (or vapor) issues from a cylindrical tube into co-flowing oxidizer [107], and counterflow flames, where a reaction zone is established between two opposing streams of fuel and oxidizer [108, 109]. The general features of the flow configurations differ between these two types of flames. A coflow flame has a two-dimensional flow field so that the species concentrations, temperatures, and flow velocities vary in both axial and radial directions. A counterflow flame, on the other hand, is quasi-one-dimensional along the normal direction of the flame, a feature that significantly facilitates the analysis of flame/sooting structures. This becomes particularly relevant if models with detailed chemical kinetics and particle dynamics are to be employed for the simulation of these flames, as the decreased dimensionality of counterflow flames
significantly reduces the computational cost. To give an example, a recent computational work [110] on an ethylene/air coflow flame using a semi-detailed chemical reaction mechanism [111, 112] and a sectional soot model was reported to take a wall time of more than 150 hours on a 400 CPU Linux cluster (i.e., 60,000 CPU hours) for convergence; similar work for one-dimensional counterflow flames typically takes no more than several hours on a PC. Also, it is generally believed that a reacting zone of premixed nature is responsible for the stabilization and attachment of the flame to the nozzle in a coflow diffusion flame. This premixed zone exists due to the inter-diffusion of fuel and oxidizer in the vicinity of the burner nozzle, where chemical reactions are inhibited by heat and radical losses to the wall [109, 113]. Therefore, coflow diffusion flames are also affected by the characteristics of this stabilizing premixed edge flame. For this reason, some fundamental aspects of diffusion combustion are thought not to be well represented by coflow diffusion flames, while counterflow diffusion flames are recognized as pure diffusion flames [109]. Finally, the residence time in coflow flames is relatively long and sometimes difficult to parameterize. In counterflow flames, residence time and stretch rate can be adjusted by varying fuel and/or oxidizer flow velocities, providing a unique way to study soot chemistry with variable residence time. Also, its much shorter residence time is more representative of the turbulent processes in practical flames [114]. Despite the above merits, it is important to point out that counterflow flames also have disadvantages. For instance, a counterflow flame is usually rather confined in space, necessitating high spatial resolution for its diagnostic. In contrast, coflow flames are more spread out spatially which makes probe sampling of the gas phase or particulates much easier.

Major differences exist in the sooting characteristics between coflow and counterflow diffusion flames. Soot formed in the fuel-rich region of a normal coflow flame is always convected downstream, towards the high temperature flame front where fuel and oxidizer are mixed stoichiometrically. As a result, oxidation of soot by oxygen and hydroxyl radical is inevitable. In fact, the widely referred *smoke point* condition is a critical condition where soot formation is balanced just by its oxidation
[104, 115-117]. But for counterflow diffusion flames, the absence or presence of soot oxidation depends on the relative position between the stagnation plane and the flame front. By adjusting the dilution ratio of the fuel and oxidizer streams (and thus the stoichiometric mixture fraction), it is possible to establish a soot formation (SF) flame where soot particles, once formed, are convected away from the flame without further oxidation [73]. In this way, the soot formation process, as well as the physicochemical features of the flame-generated soot, can be investigated with little interference from soot oxidation. Detailed descriptions of the sooting processes in diffusion flames are covered in a later section, but it can be seen from the above discussion that counterflow flames provide a well-defined, canonical configuration to complement coflow diffusion flames for soot research on diffusion flames.

The literature on soot formation is extensive and excellent reviews [18, 31, 34, 99, 107, 118-128] were periodically published to summarize research progress. Many of these reviews had a specific focus. For example, in addition to providing an excellent theoretical discussion on the general kinetic pathways of soot formation, experimental data reviewed by Wang [18] were largely based on studies of burner-stabilized premixed flames. The work of Eremin [31] focused on the formation of soot particles in shock tubes while Tree and Svesson [124] analyzed soot processes in compression ignition engines. D’Anna [125] insightfully examined both numerical and experimental aspects of nanoparticle formation in laminar flames, focusing on premixed and coflow diffusion flames, while with limited coverage on counterflow diffusion flames. Karataş and Gülder [107] offered a dedicated review on high-pressure coflow diffusion flames. Despite the importance of soot studies in laminar counterflow flames, and the many important conclusions drawn from them, there has been no dedicated review in the open literature summarizing soot research in these flames, which dates back to 1980s and continues to contribute to our expanding knowledge on soot. The special features of soot processes in counterflow flames certainly deserves such efforts. Motivated by this, we intend to provide a comprehensive survey of research outcomes resulting from investigations of soot formation
in counterflow flames, complementing existing reviews on soot research in many other flame/reactor configurations.

1.3 Structure of this review

Following is the structure of this survey of soot research in laminar counterflow flames. First, an overview of the soot formation pathways is revisited, with special attention paid to the most recent progress. The discussion follows the generally acknowledged major processes of soot formation [122]: 1) gas-phase precursor formation; 2) soot nucleation; 3) particle surface growth, particle coalescence and agglomeration; and 4) soot oxidation and fragmentation. Experimental methods for soot diagnostics are then briefly discussed to provide necessary background for following discussions on the general sooting structures of diffusion and partially-premixed flames. Similarities and differences in the soot evolution processes between coflow and counterflow flames are also highlighted. Experimental and numerical counterflow flame-based soot studies are then extensively reviewed and the materials are organized based on the various factors affecting soot formation, including fuel molecular structure, fuel additives, pressure, temperature, residence time, external fields, and flame unsteadiness. The discussion on fuel molecular structure also summarizes various quantitative indices developed to represent sooting tendencies of different fuels, which may find application in assessing and formulating surrogate fuels for sooting tendencies. These indices include the conventional smoke point, the coflow flame-based yield soot index and the newly proposed, CDF-based sooting temperature index and sooting sensitivity index. This review concludes with a brief summary and an outlook on research needs, with respect to both experimental and numerical aspects, for promoting fundamental understanding of soot formation in counterflow flames.
2. Pathways of soot formation

The physicochemical pathways from gas-phase fuels to soot particles in flames are not only complex but intellectually intriguing. It may seem impossible that structurally ordered solids can be formed under violent entropy-increasing flame conditions. In fact, as noted by Wang [18], dehydrogenation from hydrocarbon fuel molecules and the accompanied release of H\textsubscript{2} are important in decreasing the Gibbs free energy of the system, driving soot formation processes forward. Nevertheless, such a thermodynamic driving force is moderate, so the soot process is kinetically controlled and can be highly reversible [18].

Although many details are still not fully understood, decades of extensive research have led to the general belief that soot formation starts with the formation of gas-phase molecular species from fuel pyrolysis, followed by incipient ring formation, PAH growth, particle nucleation, heterogeneous surface reaction, and particle-particle interactions. For clarification, Figure 1 shows a conceptual schematic of this overall physicochemical pathway. This now well-known pathway is a frequent review subject, although new findings are added continuously. This section provides a systematic summary of the most recent discoveries in soot formation mechanisms. The material in this section will not be restricted to counterflow flames, since the fundamental chemistry involved in the process of soot formation is expected to share similarities among different flame configurations [107].

Figure 1. Schematic of overall pathways for soot formation.
It is our intention to provide a detailed description of our up-to-date understanding on soot formation. We think that such information can be a useful background knowledge to put the results of the to-be-reviewed counterflow-based fundamental soot studies into the context of understanding soot formation from first principles. Before detailed presentation, it is worthwhile to point out that the discussion below may contain historical developments that overlapped with previous reviews such as Richter and Howard [121], McEnally et al [123], D’Anna [125], Wang [18], and Karatas and Gülder [107]. Although we focused on the most recent developments after 2010, a certain degree of overlaps is unavoidable for a systematic discussion.

2.1 Gas phase precursor formation

Although polyacetylenes [129, 130] and ionic species [131] have historically been proposed as soot precursors, it is now widely accepted that the formation of PAHs is the key step in bridging the gap between gaseous fuel molecules and solid soot particles. Detailed reaction pathways are still needed to validate polyacetylenes hypothesis against measurements. The experimental results of low concentrations of chem-ions [132] and low levels of ionization for large hydrocarbons in fuel rich sooting flames [133], as well as the absence of ionic species in soot-forming pyrolysis environments [134], support evidence against the ionic mechanism. These are in contrast to the fact that the PAH hypothesis has been substantiated by numerous experimental and numerical studies [122]. For example, through a computational investigation, Stein and Fahr [135] found that the most thermodynamically stable species at high temperatures in a H2/C2H2 mixture were PAH molecules, suggesting their important roles in soot formation. The more recent work by Michelson, Violi and coworkers [136-138] suggesting the importance of non-stabilomer PAHs in soot formation will be discussed later. In addition, on a macroscopic scale, it was observed in a series of shock tube pyrolysis experiments that aromatic fuels (benzene and toluene) and cyclic compounds that can readily aromatize (e.g., 1,4-cyclohexadiene) have the highest soot yields, followed in order by species that can also have ring ruptures competing with aromatization (e.g., cyclopentadiene) and those that can
be easily fragmented (e.g., cyclohexane) [139]. Many recent PAH-based soot models with detailed chemical kinetics have gained unprecedented success in predicting the sooting characteristics of hydrocarbon flames, especially in terms of the effects of fuel molecular structures [112, 140]. Nevertheless, we are probably still not in a position to exclude the possible contribution of polyacetylenes and ionic species to soot formation, and more research is required to unravel this problem. In this study, however, we will presume that PAHs are the primary soot precursors and detailed discussions on other alternatives are beyond the scope of this review.

2.1.1 Formation of first aromatic ring

A number of reaction routes have been proposed for the growth of PAHs in hydrocarbon flames, and the relative importance of each individual pathway was shown to be notably dependent on fuel molecular structure, as well as pressure and temperature conditions [121]. For aliphatic fuels, it may be logical to expect the growth of PAHs to start from the first aromatic ring, which in many cases is benzene or phenol radical. Pioneering works on pyrolysis/oxidation of acetylene (C$_2$H$_2$) [141-143], C$_2$H$_2$/H$_2$ mixture [144], methyl chloride (CH$_3$Cl) [145], and rich premixed flames of C$_2$H$_2$ [146], ethylene (C$_2$H$_4$) [147], and 1,3-butadiene (1,3-C$_4$H$_6$) [148] suggest that the following reactions are important for benzene (A1) formation from small aliphatics:

\[
\begin{align*}
n-C_4H_3 + C_2H_2 & \rightarrow A1 - R1 \\
n-C_4H_5 + C_2H_2 & \rightarrow A1 + H \quad R2
\end{align*}
\]

where C$_2$H$_2$ is acetylene, n-C$_4$H$_3$ has the structure of HC≡C–CH=ĈH, n-C$_4$H$_5$ has H$_2$C=CH–CH=ĈH, A1- and A1 represent phenyl radical and benzene, respectively. In addition, it was shown that the addition of vinyl radical (C$_2$H$_3$) to C$_2$H$_2$ was pivotal in the production of n-C$_4$H$_3$ in flames of aliphatic fuels [148] (except for 1,3-C$_4$H$_6$ flame where H abstraction from the fuel molecule can directly lead to n-C$_4$H$_5$) and ethynyl radical (C$_2$H) addition to C$_2$H$_2$ was important for n-C$_4$H$_3$ formation [143, 146].
The importance of \(R_1\) and \(R_2\) was substantiated by Westmoreland and coworkers [149, 150], who screened various benzene formation pathways by comparing experimentally-obtained species mole fraction profiles with predicted ones in a series of low-pressure fuel-rich premixed \(\text{C}_2\text{H}_2/\text{O}_2/\text{Ar}\) flames. They employed the concept of chemical activation and considered the effects of pressure falloff, proposing rate constants of relevant reactions at different pressures instead of using high-pressure limits [149, 151]. Although the recombination of two propargyl radicals (\(\text{C}_3\text{H}_3\)) was also seen to be a promising route for benzene formation in the initial screening, they ruled out this possibility as they presumed that subsequent cyclization/hydrogen redistribution of the \(\text{C}_3\text{H}_3\)-\(\text{C}_3\text{H}_3\) adduct would result in serious difficulties [149].

Nevertheless, the role of \(\text{C}_3\text{H}_3\) in cyclization reactions has attracted considerable research effort since late 1980s. This is due partly to the recognition that \(\text{C}_3\text{H}_3\) is one of the most important resonantly stabilized radicals (RSRs) during hydrocarbon combustion. Being an RSR, \(\text{C}_3\text{H}_3\) is more stable than ordinary radicals and its chance for survival in a hostile combustion environment is greater. As a result, it can be accumulated to a much higher concentration, making its reactions attractive cyclization routes. It is also noteworthy that although \(\text{C}_3\text{H}_3\) is less reactive than ordinary radicals, the \(\text{C}_3\text{H}_3\)-\(\text{C}_3\text{H}_3\) addition is still a radical-radical reaction, so its rates can be significantly higher than those of radical-molecule reactions like \(R_1\) and \(R_2\) [152].

Miller and Melius [152] performed a numerical study on premixed fuel-rich \(\text{C}_2\text{H}_2\) flames and used a kinetic model in which the isomers of \(\text{C}_4\text{H}_3\) and \(\text{C}_4\text{H}_5\) were distinguished. The following H-atom assisted isomeric reactions were also included:

\[
\begin{align*}
n-\text{C}_4\text{H}_3 + \text{H} & \rightarrow i-\text{C}_4\text{H}_3 + \text{H} \quad \text{R3} \\
n-\text{C}_4\text{H}_5 + \text{H} & \rightarrow i-\text{C}_4\text{H}_5 + \text{H} \quad \text{R4}
\end{align*}
\]

where \(i-\text{C}_4\text{H}_3\) is resonantly stabilized between \(\text{H}_2\text{C}=\text{C}=\text{C}=\text{CH}\) and \(\text{H}_2\text{C}=\text{C}=\text{C}=\text{C}=\text{CH}\), and \(i-\text{C}_4\text{H}_5\) between \(\text{H}_2\text{C}=\text{C}=\text{CH}=\text{C}=\text{CH} \) and \(\text{H}_2\text{C}=\text{C}=\text{C}=\text{CH} = \text{CH}_2\). They argued that the benzene production rates from \(R_1\) and \(R_2\) alone were too slow to account for the experimentally-measured benzene
concentrations in the C$_2$H$_2$ flame. They showed that the rapid consumption of n-C$_4$H$_3$ and n-C$_4$H$_5$
(through R3 and R4, respectively), which were not included in earlier models, could explain the
insignificance of R1 and R2 at flame temperatures. Instead, they [152] proposed that the following
reactions are the leading route for benzene formation in C$_2$H$_2$ flames:

\[
\begin{align*}
\text{C}_3\text{H}_3 + \text{C}_3\text{H}_3 & \rightarrow \text{A1}^- + \text{H} \quad \text{R5} \\
\text{C}_3\text{H}_3 + \text{C}_3\text{H}_3 & \rightarrow \text{A1} \quad \text{R6}
\end{align*}
\]

The dominant contribution of R5 and R6 to benzene formation was supported by Stein et al. [153],
who independently modeled benzene formation rates in C$_2$H$_2$ flames. It has also been substantiated
by the following works on flames of other aliphatic fuels such as methane (CH$_4$) [154], C$_2$H$_4$ [155,
156], ethane (C$_2$H$_6$) [154], propene (C$_3$H$_6$) [157-159], propane (C$_3$H$_8$) [160], n-butane (n-C$_4$H$_{10}$)
[161], 1-pentene (1-C$_5$H$_{10}$) [162], and 1,3-pentadiene (1,3-C$_5$H$_8$) [163]. In addition, the propargyl
recombination route was also necessary to explain the relative tendency of benzene formation during
the oxidation of C$_2$H$_2$, allene (C$_3$H$_4$-A), and propyne (C$_3$H$_4$-P) in a jet stirred reactor [164]. In
particular, experimentally-observed synergistic effects [165] on the formation of PAHs and soot in
CDFs of ethylene-propane binary mixture fuels provided further evidence on the importance of C$_3$H$_3$
in aromatic species formation.

The chemical pathway for propargyl recombination is in fact more complex than it may seem in
R5 and R6. Based on computed potential energy surfaces, Miller and Klippenstein [166] found that
three varieties of energized linear C$_6$ adduct can be formed, and each can further isomerize among
various forms, dissociate into bimolecular products or back into the reactants. Additional product
channels (R7 and R8) for propargyl recombination were proposed [167] based on this theoretical
analysis, and related rate coefficients for R5 through R8 were modified to reflect more recent rate
calculations [168].

\[
\begin{align*}
\text{C}_3\text{H}_3 + \text{C}_3\text{H}_3 & \rightarrow \text{fulvene} \quad \text{R7} \\
\text{C}_3\text{H}_3 + \text{C}_3\text{H}_3 & \rightarrow \text{C}_4\text{H}_5\text{C}_2\text{H} \quad \text{R8}
\end{align*}
\]
Here, C₄H₅C₂H is 2-ethyl-1,3-butadiene and fulvene (C₅H₄CH₂) can easily be converted to benzene through H-atom-assisted isomerization [169, 170]. Of course, as pointed out by Miller and Klippenstein [166], R₅ through R₈ do not represent a complete description of C₃H₃ recombination chemistry. Nevertheless, as a lumped model, they are suitable for applications in typical flame conditions. Caution is required if they are used in low-temperature and high-pressure conditions.

Allyl radical (C₃H₅-A) is also an RSR and its combination with C₃H₃ was proposed as a potentially important cyclization step [160], especially under conditions where C₃H₅-A can be produced in large quantities such as in flames fueled by allene (H₂C=CH=CH₂) [171]. The implementation of this pathway in chemical mechanisms typically takes the following form [160]:

\[
C₃H₃ + C₃H₅-A \rightarrow \text{fulvene} + H + H \quad \text{R9}
\]

Recent developments and applications of synchrotron-based tunable VUV-ionization molecular-beam mass spectrometry, which could provide isomer-resolved intermediate species identification and quantification, have contributed significantly to our understanding on flame chemistry [172]. A number of premixed fuel-rich flames burning aliphatic fuels were investigated with this technique, and it was found that reactions involving propargyl radical (i.e., R₅ through R₉) are always important, while contributions from other pathways are somewhat fuel dependent [172, 173]. For instance, the reaction between i-C₄H₅ and C₂H₂ (R₁₀) is identified as a major benzene formation pathway (through fulvene) in both premixed [167] and diffusion [174] 1,3-butadiene flames, with comparable importance to propargyl recombination.

\[
i\text{-C}_4\text{H}_5 + \text{C}_2\text{H}_2 \rightarrow \text{fulvene} + H \quad \text{R10}
\]

Note that i-C₄H₅ can be directly formed from the dehydrogenation of the fuel molecule, and thus can be accumulated in large concentrations in 1,3-butadiene flames.

In methyl-cyclohexane (C₆H₁₁CH₃) flames/pyrolysis, stepwise dehydrogenation toward fulvene and benzene was seen to be a cyclization step of significant importance [175, 176]. This is also true for cyclohexane (C₆H₁₂) flames [177-180], where stepwise dehydrogenation leads directly to
benzene. Note that RSRs such as cyclohexenyl (c-C₆H₆) and cyclohexadienyl radicals (c-C₆H₇) are also involved in the process of cyclohexane dehydrogenation. It is also noteworthy that R₂ was shown to account for 6% of the total benzene formation in a fuel-rich cyclohexane flame; the explanation was based on specific fuel destruction pathways that can generate relatively large concentrations of n-C₄H₅ [177]. Besides, R₂ was seen to contribute to 3.5% of benzene in a fuel-rich 1-hexene (1-C₆H₁₂) flame [181].

Other benzene formation pathways include the addition of C₅ and C₁ species, notably the reaction between cyclopentadienyl (c-C₅H₅) and methyl radical (CH₃) to form benzene through fulvene [170, 182]:

\[
c-\text{C}_5\text{H}_5 + \text{CH}_3 \rightarrow \text{C}_5\text{H}_4\text{CH}_3 + \text{H} \quad \text{R11}
\]

Further hydrogen loss from C₅H₄CH₃ leads to fulvene. R11 has been tested in cyclopentene (c-C₅H₈) flames where c-C₅H₅ can easily be formed by H abstraction from fuel molecules [183]. Nevertheless, the results showed that major benzene formation pathways are still propargyl recombination and R11 only plays a minor role. Miller et al. [184] explained that at flame conditions, it is typical for c-C₅H₅ to decompose back to C₂H₂ and C₃H₃ (R12) and as a result, R₅ through R₈ become dominant aromatic ring formation routes. Interestingly, however, R11 was seen to be the dominant channel for benzene formation in a recent pyrolysis study of ethylcyclohexane (c-C₆H₁₁C₂H₃) [185].

\[
c-\text{C}_5\text{H}_5 \rightarrow \text{C}_3\text{H}_3 + \text{C}_2\text{H}_2 \quad \text{R12}
\]

The following reactions between ethynyl-substituted propargyl radical (i.e., i-C₅H₃, HC≡C–C≡C–CH₂, also an RSR) and CH₃ were considered potential contributors to benzene formation:

\[
i-\text{C}_5\text{H}_3 + \text{CH}_3 \rightarrow \text{fulvene} \quad \text{R13}
i-\text{C}_5\text{H}_3 + \text{CH}_3 \rightarrow \text{A1-} + \text{H} \quad \text{R14}
i-\text{C}_5\text{H}_3 + \text{CH}_3 \rightarrow \text{A1} \quad \text{R15}
\]

Although the presence of i-C₅H₃ has been experimentally confirmed in fuel-rich flames [186], direct evidence of the importance of R13 through R15 still awaits. In fact, based on a comparison with the propargyl recombination reactions, Miller et al. [184] argued the rates of R13 through R15 should be
very small. Also, the formation of benzene through linear C₆Hₓ molecules such as 1,3-hexadien-5-yne (HC≡CCH=CHCH=CH₂) and 1,3,5-hexatriene (CH₂=CHCH=CHCH=CH₂) were suggested to be important for premixed C₂H₄ [187] and C₂H₂ [146] flames.

From the above discussions, it is clear that—with the notable exception of R1 and R2—the majority of the important benzene formation pathways identified thus far for flames of aliphatic fuels involve RSRs, suggesting their significance.

It is useful to point out that benzene is not necessarily the first aromatic ring to form during the combustion of aliphatic fuels. There are several other aromatic species that can serve as precursors to grow to larger PAHs and soot. Bockhorn et al. [188] proposed the formation of phenylacetylene (c-C₆H₅C₂H) through consecutive addition of ethynyl radical (C₂H) and acetylene to diacetylene (C₄H₂), followed by ring closure, thus bypassing benzene. Other notable examples include the ring enlargement reaction between cyclopentadienyl radical and C₂H₂ to form cycloheptatrienyl radical (c-C₇H₈), which can then isomerize to benzyl radical (A₁CH₂⁺) [189]. This pathway R16, as shown below, was found to be important in cyclopentene flames [190].

\[
\text{R16}
\]

The recombination of cyclopentadienyl radical has been proposed as a formation pathway toward two-ring aromatics (i.e., naphthalene, A₂, C₁₀H₈) [170, 191]. Melius et al. [170] in particular suggested the following reaction paths, R17 and R18, through 9-H-fulvalenyl radical (c-C₅H₅−c-C₅H₄):

\[
\text{R17}
\]

\[
\text{R18}
\]
c-C₃H₅ + c-C₃H₅ → A₂ + 2H \textbf{R19}

Marinov et al. [161] provided rate constants for the overall reaction \textbf{R19} which were included in many kinetic models for flame simulations [192, 193]. Several other studies used the reduced pre-exponential factor for \textbf{R19} [194]. Recent investigations [195] found that agreement between experimental data and those predicted could be obtained by using a rate constant one order of magnitude slower than that suggested by Marinov et al. In addition, Mebel and Kislov [196] argued that the primary product of 9-H-fulvalenyl decomposition should be fulvalene (i.e., bicyclopentadienyldiene, c-C₅H₄=c-C₅H₄) instead of naphthalene. This motivated further theoretical study on c-C₃H₅ recombination reactions and, through a revisit of the C₁₀H₁₀ potential energy surface, Cavallotti and Polino [197] proposed an alternative reaction channel through H-azulyl radical (C₁₀H₆), which is known to decompose easily to A₂ and H atom.

Reactions involving propargyl and/or radical-substituted propargyl can also lead to the formation of aromatic species bypassing benzene. Miller et al. [184] referred the following reactions, involving methyl substituted propargyl (1-methylallenyl, H₃C–C≡C–ĈH₂), to be the most important among this class:

\begin{align*}
\text{C₃H₃ + CH₃CCCH₂} \rightarrow & \quad \text{A₁CH₂ + H} \quad \text{R20} \\
\text{C₃H₃ + CH₃CCCH₂} \rightarrow & \quad \text{A₁CH₃} \quad \text{R21} \\
\text{CH₃CCCH₂ + CH₃CCCH₂} \rightarrow & \quad \text{o-xylene} \quad \text{R22}
\end{align*}

where A₁CH₂ and A₁CH₃ represent benzyl radical and toluene, respectively and o-xylene is 1,2-xylene.

For ease of future reference, Figure 2 summarizes these important formation pathways of the first aromatic ring in flames of aliphatic fuels.
2.1.2 Molecular growth beyond the first aromatic ring

The necessity to form the first aromatic ring for the initiation of soot formation applies only to aliphatic fuels. For combustion systems fueled by aromatics, molecular growths beyond the first aromatic ring become the critical and rate-limiting steps toward soot formation. In such cases, the aromatic rings must be broken to provide the intermediate species required for subsequent molecular growth, which will lead to PAHs with increasingly complicated structures. It may not be difficult to imagine that the number of reaction pathways involved in forming these large PAH molecules grows exponentially with their sizes. Wang [18] pointed out that it is still not known whether the number can be treated as finite. On the other hand, it becomes complicated to obtain direct experimental
evidence of any proposed reaction pathways for large PAHs, caused primarily for the following reasons. First, the concentrations of large PAHs in flames are typically very low. In fact, it was reported that peak mole fractions of PAHs can drop by one order of magnitude with every growth of two fused rings [18]. Besides the slow kinetics for PAH growth, this can also be understood from the fact that as PAHs get larger, they are more likely to form soot, removing themselves from the gas phase. The result is that instruments with continuous improvements in detection limits are needed if increasingly larger species are to be measured. Second, many isomers exist for large PAHs, and isomeric-specific identification becomes progressively more difficult as PAHs grow larger. For example, as a major isomer-specific measurement technique for small molecules, the synchrotron-based photo-ionization flame-sampling molecular beam mass spectrometry (MBMS) is confronted with significant challenges in isolating isomers of large molecules due (partially) to the uncertainty of obtaining their ionization energies. Nevertheless, theoretical and experimental investigations during the past decades did provide considerable knowledge about the formation pathways of large PAHs, which is being used successfully in a variety of models for the simulation of soot.

The most well-known PAH growth pathway is the hydrogen-abstraction-C$_2$H$_2$-addition mechanism (or HACA for short) proposed by Frenklach, Wang and coworkers [141, 142, 146, 198]. In its essence, this HACA mechanism represents a repetitive reaction sequence of 1) radical activation by H-atom abstraction from stable molecules and 2) C$_2$H$_2$ molecule addition to the radical site. Taking the growth from benzene (A1) to naphthalene (A2) as an example, the HACA mechanism is realized as:

\[
\begin{align*}
R23
\end{align*}
\]

Alternatively, the addition of acetylene to phenyl radical can also lead to naphthalene formation through the following reaction:
R23 and R24 were initially included in the model for the prediction of soot formation in shock tube pyrolysis of C2H2 [141], and it was found that R23 played the dominant role. Frenklach et al. [141] attributed this to the instability of the A1CHCH adduct, which has a greater likelihood to decompose to the stable phenylacetylene molecule before further C2H2 addition can occur. After the formation of naphthalene, further hydrogen abstraction from the naphthalene molecule (by hydrogen atom) and subsequent C2H2 addition can lead to the formation of phenanthrene. Similar processes continue to form larger and larger PAHs, thus repetitive. Using the notations Ai and Ai- to represent aromatic species and its radical with i-number of rings, such a repetitive process can be represented in the following sequences:

\[
\begin{align*}
A_i + H & \rightleftharpoons A_{i-} + H_2 \quad \text{R25} \\
A_{i-} + C_2H_2 & \rightleftharpoons A_iC_2H_2 \quad \text{R26} \\
A_iC_2H_2 + C_2H_2 & \rightleftharpoons A_{i+1} \quad \text{R27} \\
or \quad A_{i-} + C_2H_2 & \rightleftharpoons A_iC_2H + H \quad \text{R28} \\
A_iC_2H + C_2H_2 & \rightleftharpoons A_{(i+1)-} \quad \text{R29}
\end{align*}
\]

It is important to note that as the related thermodynamic driving forces are only moderate, the HACA reaction sequences are highly reversible. In fact, the degree of reversibility of R25–R29 determines whether the HACA steps can contribute to PAH growth [122]. Therefore, reliable thermodynamic data of PAH species [199], from which reverse reaction rates can be calculated, become a prerequisite for accurate modelling of PAH and soot formation.

As Wang [18] pointed out, the HACA mechanism describes a process that satisfies the core thermodynamic and kinetic requirements of a sooting process. On one hand, soot precursors must
survive hostile flame environments, and in this regard, peri-condensed PAHs are known to be among the most important stable species in sooting flames. On the other hand, before further molecular growth can occur, stable PAHs must be activated to form corresponding aryl radicals, achievable via hydrogen abstraction reactions. As the most important intermediates in sooting flames, C₂H₂ serves as the building block for mass growth. In addition, these reactions have energy barriers and are highly reversible, giving temperature a critical role. When detailed structures of PAH and their radicals are considered, the addition of C₂H₂ can have many different and complicated realizations—R25 through R29 are only global representations. For more details we refer to a recent work by Frenklach and coworkers [200] in which a comprehensive list of reaction channels is provided. Despite its central importance, the HACA mechanism may be only one realization of the complex PAH growth pathways. Several other reaction sequences have also been shown to contribute to PAH growths.

Panariello et al. [201] sampled from an atmospheric fuel-rich C₂H₄/O₂ premixed flame and performed species detection using molecular beam time-of-flight (TOF) mass spectrometry, primarily targeting high molecular weight species. Subsequent fast Fourier transform (FFT) analysis of the mass spectra indicated that, in addition to C₂H₂ addition, PAH growth can be achieved with the addition of methylene (-CH₂-) into the bay region of PAHs. The relevance of odd carbon numbered PAHs with five-member rings were also inferred. In addition, through a computational investigation, Liu et al. [202] showed that the sequence of C₂H₂ addition, intramolecular hydrogen migration, CH₂ addition, and cyclization with H elimination can result in the formation of a five-member ring, as shown below.

![Diagram of PAH growth pathways](attachment:diagram.png)
Frenklach et al. [36] performed soot modeling for 1,3-butadiene shock-tube pyrolysis and found that vinyl radicals (C$_2$H$_3$) and vinyl acetylene (C$_4$H$_4$) are dominant growth agents for the formation of naphthalene, through R31 and R32, respectively. Note that both C$_2$H$_3$ and C$_4$H$_4$ are pyrolysis products of the parent fuel. Their results further suggested that dominant PAH growth pathways can be fuel dependent. Similarly, Raj et al. [203] advanced the addition of C$_4$H$_4$ to pyrene radical (C$_{16}$H$_9$-) to form benzo(e)pyrene (C$_{20}$H$_{12}$).

Shukla et al. [204] proposed the phenyl addition/cyclization (PAC) sequences for PAH growth based on their investigations of the pyrolysis products of toluene and toluene/benzene mixtures. Typically, PAC sequences begin with the addition of phenyl at a fusing site of an aromatic molecule. Further dehydrogenation and cyclization reactions lead to the formation of an aromatic species with more fused rings. Examples are given for the formation of fluoranthene (C$_{16}$H$_{10}$) from naphthalene (R33) and the growth from phenanthrene to benzo(e)-pyrene (R34) [204, 205].
In a recently proposed PAH growth mechanism, Raj et al. [203] included two pathways for phenanthrene formation initiated by phenyl addition to phenylacetylene (R35–R36), whose possibility was suggested by the mass spectra data of Shukla et al. [204].

Shukla et al. [206] further examined the mass spectra of the pyrolysis products of toluene/acetone mixtures at three different temperatures and found that mass peaks occurred in bundles, with intervals of 14-mass unit. In addition, an individual major mass peak was preceded by a peak two-mass unit (sometimes also four-mass unit) lower, which became more pronounced at higher temperatures. Based on this compelling experimental evidence and relevant chemical reasoning, they proposed the methyl addition/cyclization (MAC) mechanism as a growth pathway for PAHs, which is depicted in Figure 3 [206]. Note that the MAC mechanism explains the expansion from cyclotetrafused to cyclopentafused, and then to hexagonal networks. The sequences with blue arrows in Figure 3 provide one example. Note that the MAC pathways were also substantiated by more recent mass spectral studies of counterflow diffusion flames of C4 hydrocarbons [207, 208].
Figure 3. Methyl addition/cyclization pathways for PAH growth, shown here, are detailed pathways from benzene to naphthalene (a), naphthalene to 1H-phenalene (b) and phenanthrene to pyrene (c). Blue arrows represent methyl addition H-abstraction reactions. Numbers below each molecular structure are its molecular mass. Redrawn using data from Shukla et al. [206].

Following the pattern for biphenyl production through the addition of phenyl radical (c-C₆H₅) to benzene (R37) [36, 209], Böhm et al. [210] demonstrated more general combinatorial growth (ring-ring condensation) pathways which are primarily driven by biaryl reactions. It was also found that the ring-ring condensation pathway plays a dominant role in PAH growth during C₂H₂ and benzene pyrolysis [210].
In addition to \( \text{C}_2\text{H}_2 \), \( \text{CH}_3 \) and phenyl radicals, the resonantly stabilized \( \text{C}_3\text{H}_3 \) radical was also suggested as an important contributor to PAH growth. For instance, naphthalene can be formed by the addition of \( \text{C}_3\text{H}_3 \) on benzyl radical (c-C\( _6\text{H}_5\text{CH}_2 \)) \((\text{R38})\) [211]:

![Reaction R38](image1.png)

D’Anna and Violi [211] proposed the sequential addition of \( \text{C}_3\text{H}_3 \) to phenyl radicals, resulting in the formation of phenanthrene \((\text{R39})\).

![Reaction R39](image2.png)

Raj \textit{et al.} [212] recently recommended a PAH growth pathway from naphthalene to pyrene \((\text{R40})\) through \( \text{C}_3\text{H}_3 \) addition, and they provided kinetic parameters for each involved elementary reactions through theoretical computations based on the density functional theory (DFT). It was further suggested that analogous pathways and kinetic rates can be extrapolated to larger PAH molecules that have a zigzag site similar to naphthalene [212].

![Reaction R40](image3.png)

Aromatic radical substituted propargyl radicals were proposed to contribute to PAH growth by Stein \textit{et al.} [153]. In particular, terphenyl (i.e., diphenylbenzenes, \( \text{C}_{18}\text{H}_{14} \)) can be formed through the recombination of phenyl-substituted \( \text{C}_3\text{H}_3 \) radicals \((\text{R41})\). Note that further dehydrogenation will lead to the fused-ring structure triphenylene (\( \text{C}_{18}\text{H}_{12} \), a flat PAH consisting of four fused benzene rings).
Another important RSR, cyclopentadienyl radical (c-C₅H₅), was shown to contribute to PAH growth. Besides its self-recombination to form naphthalene (R19), its addition to indenyl radical (c-C₉H₇) can lead to the formation of phenanthrene (R42) [154]:

Indenyl radical can further contribute to pyrene formation through self-combination, or addition to benzyl radical, the global reaction of which was included in the kinetic network of Raj et al. [212].

It can be inferred from the above discussions that there exist many different pathways to PAH growth, and the relative importance of each individual pathway depends on fuel molecular structures as well as flame environments. In addition, resonantly stabilized radicals such as propargyl, substituted propargyl, cyclopentadienyl, benzyl and indenyl radicals are important growth agents. Most of these reactions are proposed only as global reactions so that further study is required for detailed theoretical analyses of the related elementary reactions to provide more accurate and pressure-dependent product channels and rate constants. On the other hand, statistical methods (i.e., kinetic Monte-Carlo) that allow PAHs to grow indefinitely by employing different classes of reactions in a jump manner [213-217], may be more practical in the context of soot simulation. This will be discussed further in Section 2.3 on soot surface growth, as recent models employ a chemical analogy [218] of soot with large PAHs, considering soot particles to be composed of stacks of PAHs and surface growth can be described by PAH planar or spatial growth at edge sites.
2.2 Soot nucleation

Soot nucleation refers to the transition of gas-phase molecular soot precursors to condensed-phase carbon materials. According to Frenklach [122] and many other researchers, this transition is probably the least known process of soot formation. Nevertheless, extensive research in the last few decades has made it convincing that high molecular weight PAHs are the dominant—if not the sole—class of molecular precursors for soot nucleation. The evidence includes transmission electron microscopy (TEM) images of soot indicating ordered structures in soot particles, believed to arise from a stacking of PAHs to form parallel layers [219]. However, definite physicochemical mechanisms explaining how the first soot particles nucleate out of PAHs under flame conditions remain research in progress.

Various conceptually different pathways for soot nucleation have been suggested [18] and are briefly discussed here. More detailed analysis can be found in the review by Wang [18]. The first pathway (with some variation in details) generally concerns the formation and growth of fullerene-like structures. In the famous graphite laser vaporization experiments by Kroto et al. [220] in which Buckminsterfullerene ($C_{60}$) was discovered, the decrease in the concentrations of other large C$_n$ clusters (except $C_{60}$) was observed with the increase in reaction residence time. The authors proposed that the consumption of these more reactive (compared to the remarkably stable $C_{60}$) large carbon clusters are due to their interactions with the abundant small carbon species to form soot. It was further argued that, at typical flame temperatures where dehydrogenation are favored, carbon tends to form stable spherical shells to satisfy the valence requirements and curved PAHs with five-member rings can contribute to the generation of the spheroidal structures [221]. Based on this reasoning, an icospiral mechanism was proposed in which the formation of soot is described as a consequence of faulty growth of fullerene. The error during fullerene formation prevents cage closure, instead it leads to further molecular growth to onion-like structures [221, 222]. Figure 4a provides a schematic for this process. The icospiral mechanism suggests fullerenic structures to be precursors for soot, however, it should be noted that although the authors wish to generalize soot formation in flame
conditions, this mechanism was based on experiments performed for a pure carbon system. Homann and coworkers [223-226] studied fullerene and soot formation in premixed flames using MBMS with photon ionization and observed that whenever soot or fullerene are formed, negative ions, covering a continuous mass spectrum up to 2,000 amu, are present. Presumably, this indicates the presence of aromers (aromatic oligomers), which were assumed to be products of PAH-PAH reactions [222, 225]. It was further argued that aromers possess many five-member rings (to explain their electron affinity) and have curved structures facilitating hydrogen loss, the features of which suggest their possible role as fullerene/soot precursors [222]. Homann explained further that once aromers are formed, the conditions of flame temperature, pressure, and intermediate species’ concentration will determine whether they continue to react with unsaturated hydrocarbons (e.g., C₂H₂) to form soot or undergo unimolecular reactions toward cage closure to form fullerene. In conventional combustion environments, where hydrocarbons are burned in air (fuel rich) at atmospheric or elevated pressures, unsaturated small hydrocarbons are abundant, favoring soot formation [222]. The pathway from aromers to soot is schematically shown in Figure 4b.

![Figure 4. Schematic of soot formation through icospiral mechanism (a), adapted from Kroto [227] with permission of the AAAS, and from aromatic oligomers (b), adapted from Homann [222] with permission of John Wiley and Sons.](image-url)
Another nucleation pathway is related to the purely chemical growth of soot nuclei from PAH species. D’Anna and coworkers [125, 215] proposed a growth mechanism in which PAHs chemically coalesce with each other to form three-dimensional, cross-linked, ring-ring aromatic structures. With continuous mass and size growth of these structure, the properties of the condensed phase can finally be acquired. This process can also be schematized by the following repetitive reaction sequences [125, 215]:

\[
\begin{align*}
A_i + H &\rightleftharpoons A_{i-} + H_2 & \text{R43} \\
A_{i-} + A_j &\rightleftharpoons A_{i+j} + H & \text{R44}
\end{align*}
\]

where \( A_i \) represents PAH species and \( A_{i-} \) their radicals and the subscript i (or j, i+j ) indicates the size of the PAHs. Note that for such a kinetic scheme—if steady state assumption is invoked for the radical species—the rate of aromatic growth can be expressed as \( k_{44}[A_i] K_{eq,43}[H]/[H_2] \), where \( k_{44} \) is the rate constant for \( \text{R44} \) and \( K_{eq,43} \) is the equilibrium constant for \( \text{R43} \). It can immediately be seen that the growth rate is dependent on the concentration ratio between H radical to H\(_2\) molecules. Therefore, it is reasonable to expect that a flame environment rich in H radicals is required in order for this chemical growth mechanism to be efficient.

In yet another pathway advocated by Frenklach and Wang [198], soot nucleation was explained to occur through physical sticking of PAHs. As the size of the aromatic species is increased beyond a certain critical value (via molecular chemical growth), they collide and begin to stick together to form PAH dimers, which continue to collide with other PAH molecules to form trimers and so on. From energy considerations, the most probable orientations for successful physical sticking are those leading to the formation of staked clusters. In this way, the PAH stacks become increasingly larger and eventually assume the properties of solid particles.

It is probably safe to say that none of the above pathways provides a universal or consistent explanation for the soot nucleation processes observed in different flame conditions. For instance, it has been argued that, from a kinetic point of view, the rates of the chemical reactions leading to
fullerenic growth are too slow to explain the observed time scale of soot inception in experiments [198, 228]. Also, experimental diffraction patterns of soot particles were reported to suggest the presence of stacked turbostratic benzenoid arrays (as opposed to the spiraling structure) [228], which were also supported by soot images taken with TEM [229]. Recent measurements on soot particle sizes in premixed flames confirm the dominance of bimodal size distributions [48, 49, 230-233], evidence against the icospiral nucleation pathway. The underlying reasoning was provided by Wang [18] as follows: The solution of the Smoluchowski coagulation equation with persistent source of nucleation shows that the particle size distribution function would quickly evolve into a unimodal if the rate of nucleation is first order with monomer concentration [18, 234], which is the case for icospiral nucleation mechanism. As a result, this disagreement with the experiments in terms of size distribution excludes the icospiral mechanism to be the dominant pathway for soot nucleation, at least for the premixed flame conditions investigated.

As previously mentioned, the PAH chemical coalescence pathway has its limitations in accounting for persistent soot nucleation in regions where the concentration of H radical is low. For molecular growth reactions to occur between PAH species (R44), one of the PAH molecules must be activated to form aryl radical, which is typically achieved through H-abstraction reaction involving H radical (R43). As a result, it is expected that in the post-flame zone of fuel-rich premixed flames (or the fuel side of non-premixed flames where H radicals exist only in very low concentrations), the rates of soot nucleation through chemical coalescence may be negligible. However, experimental data provide evidence that soot nucleation continues in such regions [230], suggesting that other nucleation mechanisms are responsible.

The primary issue concerning the PAH physical dimerization pathway is the contradiction between the need for large PAHs for dimer stability and the difficulty in forming these large PAH molecules. For the sticking of two PAH monomers to be successful in high temperature flame environments, the electrostatic and dispersive binding forces must be strong enough to counteract the entropy resistance.
for binding. This translates to the fact that the colliding PAH monomers must reach a rather large size if dimer formation is to be realized. A recent molecular dynamic study [235] confirmed that PAH dimer stability is controlled by monomer mass and solvent-accessible surface area, accompanied by the discovery that—in the case of heteromolecular dimerization—dimer stability is determined primarily by the size of the smaller monomer. Direct measurement of pyrene (four-ring PAH) dimerization rates in low temperature supersonic jets demonstrated pyrene dimerization, which is frequently modeled for nucleation, cannot proceed to a significant level at flame temperatures [236]. Similar conclusions were reached by a recent molecular dynamic study, showing that pyrene dimerization at 1500 K is almost impossible [237]. It was further estimated that a 3.6 bar partial pressure of pyrene is required for 1% pyrene to be converted to dimers at 1500 K, while in practical hydrocarbon flames, the volume fractions of pyrene are generally on the order of $10^{-6}$. In fact, based on theoretically calculated binding energy data [238-240], assuming equilibrium kinetics, a size of circumcoronene ($C_{54}H_{18}$) is required for successful dimerization at typical flame temperatures [18]. Non-equilibrium effects were also discussed by Schuetz and Frenklach [241], who performed molecular dynamics simulation of pyrene dimerization with on-the-fly quantum forces. They showed that a rovibrationally excited dimer can be formed from two pyrene molecules with a lifetime significantly longer than the collision time scale, exhibiting the possibility of further chemical reactions leading to the formation of a covalent bond between the two PAH monomers. An unresolved issue of these postulates is whether the excited dimer/initial clusters can survive constant non-reactive collisions during their lifetimes.

On the other hand, experiments have shown that peak concentrations decrease exponentially for increasingly larger PAHs. For instance, in a typical ethane counterflow diffusion flame [242] at atmospheric pressure with a benzene mole fraction on the order of $10^{-4}$, the mole fractions for the two-ring naphthalene and four-ring pyrene dropped to $10^{-5}$ and $10^{-6}$, respectively. In fact, it has been pointed out that in typical sooting premixed flames, the concentrations of PAHs decreased by roughly
one order of magnitude with the growth of every two peri-condensed rings [18]. Therefore, it is expected that the concentrations of large PAHs (i.e., $A_{19}\ C_{54}H_{18}$) will be too small to account for typical values of soot number density observed in actual sooting flames.

There is also an experimental evidence that raises questions about nucleation from pyrene. Recent aerosol mass spectrometer studies on extracted soot particles from a $C_2H_4/C_2H_2$ counterflow diffusion flame showed that species with mass 202 amu (pyrene or its isomers) contribute a very small fraction of the high-mass species [136, 137]. Even for the mass peak of 202 amu, the photo-ionization efficiency curve (PIE, discussed later in the section on experimental method) showed that it was most likely contributed by isomers of pyrene, leading to the tentative conclusion that PAH species other than the Stein stabilomers [135] may be important for soot nucleation.

The insufficiency of the above soot nucleation hypothesis in explaining experimental observations motivates the investigation of additional pathways for nascent soot formation; several research groups have made significant efforts to search for additional/alternative nucleation schemes. For example, through molecular dynamics simulations, Violi and coworkers [243, 244] showed that the aliphatic chains attached to aromatic rings may be important for facilitating the physical sticking of PAHs into clusters at high temperatures, by accommodating the excessive collision energy into the additional internal vibrational modes of these side chains.

Considering the experimental observation that, in addition to PAH clusters held together by physical forces, aromatic moieties linked by chemical bonds are also present in high temperature region of a counterflow diffusion flame [89], Mao et al. [245] pointed out the potential importance of chemical nucleation for the formation of incipient soot formation. They investigated the soot nucleation process from a pool of PAH molecules of various sizes using the reactive force-field (ReaxFF) molecular dynamic simulation, which has the capability to account for both physical and chemical reactions. Results showed that at low temperatures, even PAHs as small as naphthalene can nucleate into stacked structures via physical interactions. Nevertheless, at typical temperatures in
sooting regions of hydrocarbon flames (i.e., ~1600K), physical sticking of PAHs was not possible except for very large molecules (e.g., circumcoronene, A_{19}), which is consistent with the conclusion of Sabbah et al. [236]. With a further increase in temperature to above 2500 K, PAHs became chemically active through dehydrogenation (i.e., formation of aryl radicals). Subsequent PAH fragmentation produced polyacetylenic-like radicals or small molecules which further contributed to molecular growth. Specifically, both PAH clusters connected with a carbon bridge (i.e., a covalent bond with polyacetylenic chains) at the edge, and fullerene-like structures were observed. This study essentially supports the chemical growth pathway proposed by D’Anna and coworkers [125, 215]. However, the high temperature (> 2500K) required for H-abstraction through dehydrogenation may not fully explain the ubiquitous soot formation at much lower temperatures (~1500 K) [246, 247].

Motivated by the prediction that the zigzag edges of graphene, or finite-sized graphenes, contain radical characteristics originating from localized π-electrons, Wang [18] pointed out the potential role of aromatic π-radicals in strengthening the binding in PAH clusters. In a step forward, Zhang et al. [248] performed quantum chemistry calculations to investigate the effects of π-electrons in covalent bond formation during PAH dimerization. The bond nature (i.e., number of bonds and sites for bonding) was systematically studied for dimerization of both identical and different rectangular PAHs. Their results confirmed the possibility of covalent bonding, and that the radical character dictates the number of bonds formed, while the aromaticity of the individual six-member ring determines the bond sites. This implies that bonding is more likely to form in rings with the largest aromaticity. It must be pointed out, however, the discussion of Zhang et al. [248] was based on a zero-temperature scenario, while at combustion relevant temperatures the dimers formed were found to be thermodynamically unstable [249], primarily due to the large decrease of dimerization entropy. Nevertheless, the authors considered that the lifetime of dimers may be longer than the molecular collision time scale so that they, once formed, could still contribute to the nucleation of soot particles.
It remains to be proved whether these unstable dimers can survive constant collisions from molecules of other combustion gases.

The role of cation-π in enhancing the stability of PAH clusters, inspired by previous astrophysical studies in which the existence of PAHs in the interstellar medium [250-252] and the enhanced PAH binding by a metal cation [253, 254] were discussed, has recently drawn attention from researchers on soot. Recognizing that the most abundant cations in hydrocarbon flames are \( \text{CHO}^+ \), \( \text{C}_3\text{H}_5^+ \) and \( \text{H}_3\text{O}^+ \) resulting from chemi-ionization [255, 256], Chen and Wang [257] performed \textit{ab initio} and density functional theory calculations to investigate the binding energies and structures of complexes formed from the association of these chemi-ions and 1 to 4 ring PAHs. Their results showed that although the cation-π interactions make a positive contribution, the cation-enhanced bindings are still too weak to account for soot nucleation from PAH clusters. However, similar studies on the interactions between cations of transition metals and benzene molecules [258, 259] have shown very strong binding energies among the complexes \( \text{M}^{z+}(\text{Bz})_n \), where \( \text{M}^{z+} \) is the metal cation (i.e., \( \text{Fe}^{2+} \), \( \text{Co}^{2+} \), \( \text{Cu}^{2+} \)), \( \text{Bz} \) refers to benzene, and \( n \) is the number of benzene molecules in the complex. Such enhanced binding strength indicates the potential of \( \text{M}^{z+}(\text{Bz})_n \) to form the first soot nucleus [258]. This is especially relevant considering many trace metals present in practical hydrocarbon fuels, as well as in engine lubricating oils. However, it must be pointed out that although theoretically plausible, direct experimental evidence has yet to be produced for the facilitation of \( \text{M}^{z+}(\text{Bz})_n \) in soot nucleation, despite the fact that the effect of metal addition in soot formation was studied decades ago [260]. The problem is complicated because, besides its potential role in nucleation, a metal additive can also catalytically reduce the formation of molecular soot precursors [261, 262], enhance the oxidation of soot once formed [45, 261, 263], and suppress particle-particle coagulation due to repulsive forces [260, 264].

More recently, Kraft and coworkers [265, 266] noticed that curved PAHs (cPAH), with pentagonal ring integrated in their structures, are featured with significant molecular flexoelectric dipole
moments. They further investigated the possibility of dipole-induced homodimerization of cPAHs, only to find that the binding energy between cPAHs is comparable with or even lower than that of planar PAHs [267]. This is because the steric effects of cPAHs caused a reduction in dispersion contribution, and thus cancelled the effects of dipole-dipole interactions [267]. Nevertheless, it was shown that cPAHs can bind strongly with chem-ions, the interactions between which are long-ranged and can be significant especially in combustion systems where concentrations of relevant species are low [268]. The proposal that the interaction between cPAHs and ions may play an important role in soot formation [266, 268] was also supported by recent experimental results that demonstrated enhanced physical clustering with cations [54, 269]. Nevertheless, further theoretical and experimental efforts are needed to confirm this ion-induced nucleation mechanism.

From a modelling perspective, soot nucleation must be quantitatively accounted for in any predictive soot models because it bridges the treatment of gas phase and surface reactions, in addition to serving as a turning point beyond which particle dynamics is initiated. Historical treatments are largely empirical and include the one-step nucleation from C₂H₂ [270], as shown in R45, and that from benzene in R46 [271]. Many modern soot inception models follow the work of Frenklach and Wang [198] and describe soot nucleation with irreversible [198] or reversible dimerization [272] of peri-condensed aromatic species, most notably pyrene molecules. Although these models successfully explain many experimental observations, there has been compelling evidence suggesting that these dimers, linked by physical forces, are not stable enough at flame temperature to accommodate further growth [243], and that chemical bonds between monomers are essential for the formation of stable PAH clusters. The main challenges for current chemical nucleation mechanisms, mentioned above, are that they involve reactions with high-energy barriers (leading to slow reaction rates), which are not perfectly compatible with certain experimental observations. One example is the experimentally-observed persistent soot nucleation in the post-flame zone of rich premixed flames, which is characterized by low temperature and low concentrations of reactive radicals (i.e., H atom).
Recently, a new nucleation mechanism, termed *clustering of hydrocarbons by radical-chain reactions* (CHCR) was proposed by Johansson *et al.* [138]. The CHCR mechanism highlights the importance of chain propagation of resonantly stabilized radicals (RSR). In particular, the molecular growth process initiates with repetitive additions of \( \text{C}_2\text{H}_2 \) or vinyl radical (\( \text{C}_2\text{H}_3 \)) to RSRs (e.g., \( \text{C}_3\text{H}_3 \) or \( \text{c-C}_5\text{H}_5 \)), generating new RSRs with increasingly higher molecular weights. These RSRs can subsequently cluster with different types of hydrocarbons, including stable PAHs and unsaturated aliphatic species, while preserving the radical nature. As such, molecular growth can proceed without the need for high energy barrier activation reaction, leading finally to covalently-bonded large clusters (incipient particles). In this way the CHCR mechanism avoids the challenges for existing chemical nucleation mechanisms. It also overcomes the problem of insufficient concentration of large PAHs because RSRs can cluster with a wide range of different hydrocarbons [138]. The CHCR mechanism was motivated partially by experimental observations (through VUV-MS measurements) of a class of increasingly larger RSRs in sooting flames. Note that the chain propagation reactions involving RSRs has been confirmed by quantum computations [138]. However, detailed rates for the various proposed radical chain propagation reactions still need to be determined so that a complete mechanism can be assembled to test the capabilities of the CHCR mechanism in predicting the formation of large molecular species and soot [273].

### 2.3 Soot surface growth, particle coalescence, and agglomeration

Surface growth refers to the process in which gaseous species attach to the surface of pre-existing soot particles, and their subsequent incorporation into the particulate phase [119, 274]. It is generally believed that although inception is the first step for soot formation, providing the nucleus, the majority of soot mass comes from surface growth [274]. For instance, the results of a numerical simulation study on soot formation in ethylene/air coflow diffusion flames [72] showed that the radially
integrated soot mass growth rate from surface reactions were more than an order of magnitude higher than from new particle inception. Similarly, Kennedy [275] showed that more than 90% of the final soot volume is produced from surface growth in counterflow diffusion flames.

Although detailed physicochemical processes of surface growth are still not fully understood, semi-empirical descriptions of its rates were proposed decades ago, initially for premixed flames [274, 276], and later for diffusion flames as well [277-279]. These early studies generally showed a first-order kinetics with the gas phase soot growth agent, for which acetylene was considered to be a primary candidate [276]. The rate of the mass adding reaction \( \text{C}_2\text{H}_2 + n\text{C}_{\text{soot}} = (n+2)\text{C}_{\text{soot}} + \text{H}_2 \) then took the simple form of \( \omega_g = k(T)f(S)[\text{C}_2\text{H}_2] \), where \( \omega_g \) is the soot growth rate, \( k(T) \) is the rate constant dependent on temperature, \( T \), and \( f(S) \) is a function of soot surface area [270]. The authors argued that because active sites may decrease in concentration with the extent of soot growth due to surface aging [119, 129], \( \omega_g \) should not be directly proportional to \( S \). They went on to propose, \textit{ad hoc}, that the number of active sites is proportional to the square root of \( S \), and the soot growth rate can be written as:

\[
\omega_g = k(T)[\text{C}_2\text{H}_2]\sqrt{\pi \left( \frac{6M_c}{\pi \rho_c} \right)^{2/3} [\text{C}]^{1/3} (\rho N)^{1/6}} \quad \text{Eq. 1}
\]

where \( M_c \) is the molar weight of carbon, \( [\text{C}] \) is the molar concentration of solid carbon, \( N \) is the number of particles per unit mixture mass, and \( \rho_c \) is the density of soot [270]. Later experimental determination of mass growth rates following streamlines in coflow diffusion flames indicated that \( \omega_g \) was dependent on acetylene concentration raised to a power of \( n \), slightly larger than unity [277, 279]. This difference, as compared to \text{Eq. 1}, was thought to be caused by soot oxidation (discussed in detail in a subsequent section) which occurred during the experiment, thereby masking the net soot growth rate [279]. After accounting for soot oxidation, \( n \) was found to be close to unity [279].

More physically based description of surface growth was stimulated by the hypothesis of chemical similarity between soot particles and large PAHs that provided a possibility to treat surface reactions through elemental reactions [122]. In this regard, the surface analogy of the gas phase HACA
mechanism—that is the surface-HACA mechanism—was proposed to describe soot mass growth through the following reaction sequences [198, 280]:

\[
\begin{align*}
\text{C} \text{soot} - \text{H} + \text{H} & \rightarrow \text{C} \text{soot}^{'} + \text{H}_2 & \text{R47} \\
\text{C} \text{soot}^{'} + \text{H} & \rightarrow \text{C} \text{soot} - \text{H} & \text{R48} \\
\text{C} \text{soot}^{'} + \text{C}_2\text{H}_2 & \rightarrow \text{C} \text{soot} - \text{H} + \text{H} & \text{R49}
\end{align*}
\]

where \( \text{C} \text{soot} \) is bulk soot and \( \text{C} \text{soot} - \text{H} \) represents a saturated armchair (discussed later) surface site, which needs to be activated (via hydrogen abstraction) from the aryl radical site \( \text{C} \text{soot}^{'} \) before a \( \text{C}_2\text{H}_2 \) molecular can be added to increase the mass/size of the bulk carbon particle. The rate of hydrogen abstraction reaction (R47) can be obtained by kinetic analogy of similar reactions with aromatic species (e.g., benzene).

In addition to H radical, OH [146, 280], CH\(_3\), C\(_2\)H, C\(_3\)H\(_3\), and other hydrocarbon radicals, may also initiate hydrogen abstraction reactions. Hwang and Chung [281] proposed and tested a modified surface-HACA growth mechanism, emphasizing the roles of CH\(_3\), C\(_2\)H, C\(_3\)H\(_3\) in the surface site activation reactions to explain measured soot growth rates in CDFs, where H radical concentration was relatively low, for example in the relatively low temperature soot growth region in SF counterflow diffusion flames.

A parameter \( \alpha \), having a physical representation of the fraction of surface sites available for chemical reactions, was introduced by Frenklach and Wang [198] to account for the surface aging of soot particles. In principle, the extent of soot aging, and thus the value of \( \alpha (\leq 1) \), should depend on both particle size and soot surface temperature [187]. By testing several different premixed flames, an empirical correlation of \( \alpha \) was proposed:

\[
\alpha = \tanh \left( \frac{a}{\log \mu_1} + b \right) \quad \text{Eq. 2}
\]

where \( \mu_1 \) is the first moment of soot size distribution function and \( a \) and \( b \) are linear functions of temperature, which could be determined by matching with experiments. These expressions were developed based on soot measurement data for premixed flames; and there are other relations that
were used for different flame configurations [282, 283]. The Arrhenius type expression of $\alpha = \min (c \exp (d/T), 1)$, in particular, was frequently used in coflow diffusion flames [71, 284], where $c$ and $d$ are fitting constants. The value of $\alpha$ was also sometimes assumed to be a constant, for example in a previous counterflow flame simulation [78], unity value was used and a much smaller constant of 0.078 was employed by Dworkin et al. [282]. Note also that in the surface-HACA growth mechanism, the C$_2$H$_2$ addition step (R49) was assumed to be irreversible, motivated by the fact that the dependence of thermal decomposition of the surface intermediates on flame temperature would be overpredicted if reversible reaction was included [122, 198]. Nevertheless, the complete exclusion of reversibility then underestimated the temperature dependence observed experimentally and the inclusion of $\alpha$ can in a way be considered a remedy [122]. A theoretical study was recently performed [285] to microscopically investigate the surface properties of equilibrated pyrene and coronene clusters [239, 286] (as a representation of nascent soot particles), demonstrating a decreasing (increasing) trend of surface availability with increased particle size (temperature). Note that the relationship with temperature is different from the common concept of a decreasing $\alpha$ with the increase in temperature; the authors explained this difference by pointing out that surface availability is a local property, while $\alpha$ is a global property of the soot ensemble. An upper limit of 0.8–0.9 for $\alpha$ was also proposed [285], considering the pocket surface is inaccessible for gaseous species because of their shorter life time as compared to surface reaction time.

Besides C$_2$H$_2$ addition, the importance of PAH addition on particle surface in soot mass growth was also demonstrated [272, 287-289]. A modelling study on soot formation in a high-pressure coflow diffusion flame [72] showed that at pressures higher than 6 bar, the soot growth rate through PAH addition can even overtake the surface-HACA growth rate. D’Anna et al. [290] also showed that a physical mechanism such as PAH addition is more likely to contribute to nanoparticle formation and growth in the pyrolytic region of soot formation in CDF, where temperatures and H radical concentration were relatively low.
From a first principle, the types and rates of reactions on soot surfaces may depend on many parameters including temperature, size, morphology, detailed chemical composition, type, and steric confinements of reactive sites on the particles (note that the original surface-HACA mechanism [198] considered only the armchair site, and site density was pre-specified, while surface reactions on other types of sites may also be possible [291]). In addition, more recent experimental investigations in premixed flames provide both direct (aerosol mass spectrometry, Fourier-transform infrared (FTIR) and Raman spectroscopy) and indirect (TEM and atomic force microscope (AFM) [97]) evidence for the existence of aliphatic chains covering the aromatic core of nascent soot particles. These findings necessitate the description of mass growth reactions on different surface sites so that the chemical and structural details of soot particles must be tracked in soot models [18]. Toward this goal, and considering the chemical analogy of soot with PAHs (as pioneered by Frenklach [218]), Singh and Frenklach [292] and Kraft and coworkers [213, 216, 293] differentiated soot surface sites (as shown in Figure 5) and developed detailed surface mechanisms that take into consideration the different reactions on the various sites. In addition to conventional H abstraction and C\textsubscript{2}H\textsubscript{2} addition reactions in an armchair site, as discussed in [280], surface reaction can also include the formation of a five-member ring by the addition of C\textsubscript{2}H\textsubscript{2} at zigzag sites and its desorption [294], free-edge ring growth (via C\textsubscript{2}H\textsubscript{2}) and desorption [218], five- to six-member ring, as well as six- to five-member ring conversion [291, 295, 296], five- and six-member bay closure [295, 297], five-member ring migration [291, 295], benzene addition to form bi-phenyl-like structures and so on. More details of the many surface reaction classes can be found in Refs. [213, 216, 298, 299]. These reactions can be followed in detail for a single seed PAH molecule (e.g., pyrene) using a Monte Carlo algorithm, resulting in the kinetic Monte Carlo aromatic site (KMC-ARS) model [216], which can specifically track the shape and chemical identity of the growing PAH. Note that soot particles were modeled by Kraft and coworkers [216, 300] to be composed of *planar* PAHs, which contain bulk carbon atoms and edge carbon atoms. It is on these edge atoms (i.e., soot sites) that surface reaction can take place.
Fig. 5. Typical PAH structure showing different types of surface sites. *Five* indicates that the site involves a five-member ring. Adapted from Singh *et al.* [292] with permission of Elsevier.

The 3-D growth of PAH structures was also studied by Frenklach and coworkers [292, 299, 301, 302] and Violi and coworkers [303-305], partially motivated by the fact that the originally planar PAH structure can develop curvatures by incorporating five-member rings [228]. Evidence was provided by high resolution TEM (HRTEM) images which showed curved PAH lattices [306, 307]. The presence of curvature in PAHs have implications in terms of soot inception, especially considering the potential interaction (yet to be explored) between polar curved PAHs and chemi-ions [308]. Particles consisting of curved PAH lattices are also shown to be more easily oxidized [309], and a recent soot growth model by Yapp *et al.* [310] managed to estimate PAH curvature with information about the number of five- and six-member rings in the PAH structure.

Although surface growth accounts for the majority of soot mass, ultimate particle size and morphology are largely influenced by particle-particle interactions (i.e., coagulation). Collisonal interactions between particles can be generally classified into coalescence and agglomeration. Coalescence typically indicates the disappearance of the boundary between two particles in contact, resulting in a reduction of the total surface area; while agglomeration refers to the sticking of particles with point contact (conserving surface area). TEM images of nascent particles shows a spherical shape; while mature soot particles generally have nearly spherical primary particles collected into
fractal aggregates with varying numbers of primary particles [311, 312]. The spherical nature of the nascent/primary particles is probably a major motivation for an early model [198] to treat particle coagulation as purely coalescent, i.e., colliding particles merging to form a larger spherical particle. However, later investigations [313-315] showed that sphericity can also be attained by point-contact particle aggregation with simultaneous surface growth. Under conditions where colliding partners were mostly small particles (in regions where soot inception rates were high and newly incepted particles abundant), surface growth effectively smoothed out the aggregates towards highly spherical particles. On the other hand, in environments where colliding partners were large (e.g. in the post-flame zone of rich premixed flames), the smoothing effect of surface growth became relatively less important and a chain-like fractal was more likely to result.

Nevertheless, coalescent growth is still highly likely for nascent particles with liquid-like behavior. There is also experimental evidence that under high temperature conditions (> 1500K), even mature chain-like soot aggregate can experience in-particle sintering, and thus reduction of surface area and increased primary particle size [316]. In a recent detailed soot model [317], sintering was considered to be quasi-instantaneous for primary particles with a size smaller than a critical value ($D_{pp, crit}$), while for larger particles, an empirical model was used, with characteristic sintering time between two primary particles having an Arrhenius-type expression.

Complicated surface growth and particle-particle interactions would certainly result in polydispersed soot ensembles with complex particle size distributions (PSD). Recent research in high-fidelity measurement of PSD using differential mobility analysis (DMA) (see e.g., [49, 52, 54, 97, 318-321], with a detailed discussion provided by Wang [18]) opened new possibilities for understanding the above-described processes through an inverse problem [18, 50, 322, 323]. The PAH-PP soot model proposed by Kraft and coworkers [300, 317] had several parameters, such as soot density, smoothing factor, pre-exponential factor, activation energy of sintering, and fractions of reactive sites, that were optimized by comparison against experimental PSDs. Through analysis of

48
the PSD data obtained from AFM and scattering measurements, D’Alessio et al. [324] found that the sticking coefficient of coagulation (i.e., the actual measured coagulation rate divided by the theoretical value obtained from gas kinetic theory) for the nascent nanoscale organic carbon particle (NOC, <3 nm) was orders of magnitude smaller than the larger soot particles. Additional PSD measurements using both DMA and AFM confirmed that small particles have less propensity to coagulate at flame temperatures [325, 326]. By comparing the measured and computed PSD, Sirignano and D’Anna [219] further demonstrated that coagulation efficiency depends on both particle size and temperature. Small particles had low coagulation efficiency even at room temperature and they decreased further with the increase in temperature. On the other hand, coagulation efficiency for large particles did not have strong temperature sensitivity and exhibited a high value for the entire temperature range examined.

2.4 Soot oxidation and fragmentation

Soot oxidation is a process that counteracts soot growth and it is also the primary mechanism for their removal from combustion exhausts. Therefore, a mechanistic understanding of soot oxidation is of both fundamental and practical importance. A comprehensive review of early experimental and kinetic studies on soot oxidation (before 2001) was provided by Stanmore et al. [327] and a recent informative account by Frenklach et al. [200]. It is generally recognized that the most important soot oxidants are OH radical and molecular O₂, with contributions also from O atom, H₂O, CO₂ and NO₂ [327, 328]. Note that NO₂ can be particularly important at low temperatures under conditions like the regeneration of a diesel particulate filter (DPF) [329, 330].

At high temperature flame conditions, OH radicals [331-334] are particularly effective in oxidizing soot, and this can be especially true in CDFs, where concentrations of molecular O₂ in the sooting zone are relatively low. From a modeling perspective, soot oxidation by OH was typically described by the collision of a soot surface site (C_{soot-H}) with OH [198, 280], subject to collision efficiency that can be determined experimentally [334-337]. A widely-used value of this collision efficiency is 0.13,
as obtained by Neoh et al. [334] as a lower bound value. D’Anna and coworkers [214, 215, 338, 339] also considered an activation energy for such reactions, which was estimated from similar reactions for benzene and PAHs. Although such relatively simple models have been successful in predicting experimental flame data, recent theoretical studies [340-342] found that the reaction of OH + C_{soot-H} (using benzene as a prototype) does not lead to CO expulsion, and thus was not considered to be a relevant soot oxidation pathway. Instead, OH reacting with armchair (oxidation more likely at armchair site [343, 344]) soot surface radical site C_{soot} was shown to form various oxy-radicals, which then decomposed to eliminate CO. Therefore, in a most recent soot oxidation study [200], soot oxidation by OH was considered to occur only through its reaction with reactive radical surface sites.

Soot oxidation was recently studied in an environmental TEM [345]. Although the feeding gaseous reactant was molecular O_2, the high-energy electron beam of the TEM was seen to be very effective in ionizing O_2 molecules. It was thus expected that the active soot oxidants were in fact radicals such as O atom or ion. The results confirmed that a d^2-law also holds for surface oxidation of primary soot particles, regardless of their location inside the soot aggregate. The study also provided clear evidence of both the internal and surface burning modes of soot particles, confirming previous findings using conventional TEM images [346-348]. For mature particles with a thick graphite shell [349], surface oxidation dominates as it would be difficult for oxygen to diffuse across the outer shell into the core of the particles. These experimental results are believed to be helpful to model developments for soot oxidation.

Soot oxidation by molecular O_2 has recently attracted significant research attention [292, 345, 350-361], motivated at least partially by the issue of DPF regeneration. Early models described soot oxidation by O_2 as a single step: C_{soot} + O_2 → products, with rate coefficients taken by chemical analogy to the reaction of phenyl radical with O_2 [280], later updated using rates calculated for O_2 oxidation of pyrene radical (4-pyrenyl) to form phenanthrene radical (4-phenanthryl) [293]. However, this updated rate [293] was noted to be slow [362] so that additional detailed mechanistic studies were
performed and results analyzed to identify more favorable oxidation pathways [351, 363, 364]. Sendt and Haynes [365-367] investigated O₂ addition on different types of surface sites, providing understanding of the mechanism on O₂–surface sites interactions. In their studies, however, multiple adjacent radical sites were involved, which may exist only in exceedingly small concentrations considering the fraction of radical sites among surface sites were already low (10⁻² to 10⁻⁴). In this regard, Raj and coworkers used quantum calculations to re-examine reaction pathways of O₂ oxidation at free-edge [362] and zig-zag edge sites [368]. Only one radical site was present in the model reactant, and the computed reaction pathways and high-pressure limit rate constants were provided for potential incorporation into detailed soot models.

Two prominent phenomena with respect to soot burnout in oxygen-containing environments have been observed experimentally: 1) Particles containing curved PAH structures oxidize notably faster than those composed of planar PAHs [309, 359, 369] and 2) Soot oxidation reactivity tends to decrease as oxidation progresses [370]. Interestingly enough, as explained by Singh and Frenklach [292], both these enhanced and reduced soot oxidation reactivity can be explained by the presence of five-member rings in soot structures. On one hand, quantum calculations showed that activation energy for the oxidation of curved PAH (represented by 1-corannulenyl radical with a five-member ring in the center) is notably lower than that of planar PAH (represented by 4-pyrenyl radical) [359]. Sterically-resolved kinetic Monte Carlo simulation on large graphene sheets also confirmed that the oxidation rate of the curved PAH substrate was faster than planar ones, especially during the initial stage of oxidation [200]. The underlying mechanism was further identified: Curved PAH substrate tends to form a larger number of sites with edge five-member rings that can potentially desorb to expel carbon into the gas phase (i.e., not due to actual oxidation reactions) [292]. On the other hand, as oxidation progresses, many five-member rings at armchair sites accumulated, due to oxyradical decomposition. These five-member rings then formed zig-zag edges, which tends to decrease soot reactivity [292, 350]. To summarize, the detailed work of Singh and Frenklach [292] confirmed that
the effects of the five-member ring on soot oxidation depends on their position in the graphene substrate, emphasizing the importance of sterically-resolved simulations.

In addition to the reaction between oxygen and radical surface sites, the possibility of O$_2$ reacting with saturated C$_{\text{soot}}$-H sites was also investigated [200]. It was found that the possible reaction channel is for O$_2$ to abstract the H radical, forming HO$_2$. Nevertheless, the activation energy of such a reaction is high, making it less favorable than the oxidation at radical sites [200].

Soot fragmentation can be thought of as a reverse process of coalescence and agglomeration. Fragmentation can occur during thermal treatment of aggregated particles in an inert environment [371], but are more likely to happen during the oxidation processes. As discussed by Sirignano et al. [372], two types of fragmentation are possible: The first has to do with the fragmentation of large aggregates into smaller ones, or into isolated primary particles (i.e., aggregate fragmentation, reverse process of agglomeration); The second type is the fragmentation of primary particles into even smaller ones (i.e., particle fragmentation, reverse process of coalescence).

*Aggregate* fragmentation occurs when oxidative material are removed at the contact points linking primary particles; *particle* fragmentation is related to the internal structure of an individual particle [372]. Evidences for both types have been observed in different configurations such as two-stage burners [355-358], coflow [373], and counterflow diffusion flames [372]. Visual tracking of a particular single aggregate in an environmental TEM provided direct confirmation that the thickness of the bridge connecting primary particles is continuously reduced by oxidation before breakup, releasing two smaller aggregates [345]. These results show that fragmentation is an integral process of soot evolution and its accurate description is important if particle morphology, average sizes—and especially size distributions [374]–are to be accurately predicted for soot particles experiencing oxidation.

Due to the scope of the present review, oxidation of soot by NOx, H$_2$O and CO$_2$ will not be discussed further; interested readers may find relevant information elsewhere [329]. It is, however,
relevant to note, as Frenklach et al. mentioned in a recent publication [200], that soot oxidation chemistry is very rich and could be even more complicated than soot growth. As such, it is clear that more experimental and numerical work is needed for the development of predictive models of soot oxidation (and associated fragmentation) which will account for sterically-resolved surface reactions, effects of particle nanostructures, and direct coupling of the surface and gas phase environments.

2.5 Models of soot formation

With the above subsection providing physicochemical descriptions of soot formation processes, we intend to present an overview in this subsection on how these processes are numerically modeled. In fact, some modelling perspective has already been given, but here we will focus more on the numerical techniques used to deal with particle dynamics. Historical developments of soot models can be found in the review by Kennedy [120].

In many recent models for soot formation, soot inception was modelled as the physical collision of large PAHs such as pyrene [282, 375-377], or an ensemble of larger aromatics up to coronene [111, 140, 378]. In these studies, nucleation reactions were generally assumed to be irreversible with zero activation energy [198]. Recently, reversibility of the inception process was considered [288, 289, 379], and statistical mechanics were employed to determine the enthalpy and entropy of the PAH dimerization processes, through which equilibrium constants and reverse rate coefficients for soot nucleation reactions can be evaluated. The growth from gas-phase to nanoparticles was modeled as a purely chemical growth process in Refs. [136, 137, 217, 304, 380, 381]. The relevant model such as Atomistic Model for Particle Inception (AMPI) model [304, 380] and Stochastic Nanoparticle Simulator (SNAPS) [136, 137, 217, 381] were shown to correctly predict the order of magnitude of the nascent particle concentration.
Once incepted, nascent soot particles will experience surface reactions and particle-particle interactions, and their sizes and morphology will change. For the model of these processes, a particle population balance equation needs to be solved, which can be written generally as:

\[
\frac{dN_i}{dt} = \frac{1}{2} \sum_{j=1}^{i-1} \beta (j, i - j) N_j N_{i-j} - \sum_{j=1}^{\infty} \beta (i, j) N_i N_j
\]

where \(N_i\) is the number density of particle of size \(i\). \(\beta (i, j)\) is the collision kernel between particles with sizes \(i\) and \(j\). The first two terms on the right-hand side are respectively the source and sink terms due to coagulation process. Note the specific form of the collision frequency depends on the coagulation regime (free-molecular, continuum, and transition), which is characterized by the Knudsen number, \(K_n = 2 \lambda_f / d\), where \(\lambda_f\) is the mean free path for the gas-phase and \(d\) is the particle diameter. \(\beta (i, j)\) is a non-linear function of the sizes of the colliding particles, and it is quite difficult to find a closed-form solution for Eq. 3. In this regard, several approximate numerical techniques such as Monte Carlo (MC) method, method of moments and discrete sectional method have been proposed.

Monte Carlo (MC) based models [382-384] could provide accurate results by tracking the evolution of number density through an assemble of particles governed by stochastic processes [382]. Nevertheless, these simulations are computationally demanding so that their applicability has so far been limited to simple configurations such as zero-dimensional reactors [385]. In contrast, the other deterministic models including method of moments and sectional method have been successfully applied to more challenging multi-dimensional environments.

For the method of moments (MoM), the detailed description of particle dynamics (i.e. Eq. 3) is reformulated in terms of the moments of particle size distribution function (PSDF). Specifically, the \(r^{th}\) moment \(M_r\) is defined as [386]:

\[
M_r = \sum_{i=1}^{\infty} m_i^r N_i.
\]

where \(m_i\) is the mass of particle of size class \(i\). By taking the moment at both sides, Eq. 3 can be rewritten as [386]:

54
\[
\frac{dM_0}{dt} = -\frac{1}{2} \sum_{i=1}^{\infty} \sum_{j=1}^{\infty} \beta(i,j) N_i N_j \\
\frac{dM_1}{dt} = 0 \\
\frac{dM_r}{dt} = \frac{1}{2} \sum_{k=1}^{r-1} \binom{r}{k} \left( \sum_{i=1}^{\infty} \sum_{j=1}^{\infty} m_i^k m_j^{r-k} \beta(i,j) N_i N_j \right) \\
r = 2, 3, \ldots
\]

Eq. 5

Eq. 6

Eq. 7

Numerical scheme such as the interpolative closure or variants of the quadrature method (see Ref. [375] and the references therein) is necessary to close the system of equations for the moments. Following the work of Guo et al. [387], the one-dimensional transport equations for the \( r \)th moment can be written as:

\[
\frac{\partial M_r}{\partial t} + \frac{\partial}{\partial x} (u M_r) = \frac{\partial}{\partial x} \left( D_{p,1} \frac{\partial M_{r-2/3}}{\partial x} \right) - \frac{\partial}{\partial x} (V_T M_r) + \dot{M}_r
\]

where \( x \) denotes the Cartesian coordinate, \( \rho \) and \( u \) are the density and gas velocity, respectively. \( V_T \) is the thermophoretic velocity of soot particles, \( D_{p,1} \) is the diffusion coefficient for the smallest soot particle. \( M_{r-2/3} \) is the fractional moments obtained by interpolation between integer moments. \( \dot{M}_r \) is the source terms due to nucleation, coagulation, surface reaction (including oxidation), and PAH addition. Similar to the coagulation term, the other source terms are also needed to be rewritten in terms of soot moments[386], and they generally involve fractional moments.

In principle, a knowledge of all these moments is equivalent to a knowledge of PSDF itself [386]. However, in most cases the interested soot properties can be adequately represented by a finite number of moments (e.g., the first three or six moments). For example, the overall number of particles can be represented by the zeroth moment, \( M_0 \); the total volume of particles is the first moment \( M_1 \) (i.e. soot mass) divided by soot particle density; the average diameters of the particle ensemble can be derived from the ratio of for example, \( M_1/M_0 \). As a result, the method of moments could provide many expected soot properties, and in the meanwhile, keep the computational cost at a relatively low level. Therefore, it has been widely employed to compute soot particle dynamics of both laminar [122, 140, 198, 280, 375, 386, 388-391] and turbulent flames[392-394].

55
Discrete sectional method [56, 214, 282, 338, 376, 379, 395-400] is another widely used method to approximate the population balance equation. In sectional method, the continuous size distribution was discretized by a finite number of sections (or bins). Each section has a representative particle size (or mass) and the mass/volume ratio of two consecutive sections is described by a constant spacing factor. It is obvious that the PSD can be resolved more accurately if more sections are used. In reality, the choice of section number depends on the tradeoff between the approximation accuracy and computational cost.

The solution for population balance equation with the sectional method is straightforward, since the prescribed sizes $i$ and $j$ directly correspond to representative particles for the sections. Similarly, the one-dimensional transport equation for the particle number density ($N_i$) in each section can be derived as:

$$\rho \frac{\partial N_i}{\partial t} + \frac{\partial}{\partial x}(\rho u N_i) = \frac{\partial}{\partial x} \left( \rho D_{soot,i} \frac{\partial N_i}{\partial x} \right) - \frac{\partial}{\partial x} \left( \rho N_i V_T \right) + \rho \omega_{soot,i}$$

Eq. 9

where $D_{soot,i}$ is the diffusion coefficient of soot particle in section $i$. Note the Brownian diffusion of soot particles is relatively weak and in some cases its effects on particle transport can be neglected [401]. $\omega_{soot,i}$ is the soot source term for section $i$ including nucleation, coagulation, and heterogeneous soot surface processes. Incipient soot particles due to nucleation reside in the first few sections. Through coagulation or surface growth, particles in lower section can move to higher sections. On the other hand, particles in higher sections can move to lower sections by oxidation.

It is noted that Eq. 9 assumes that all soot particles are spherical (i.e., without considering the aggregate structures). This assumption was preferred in several soot models [56, 214, 378, 379, 397, 398] due partly to its low computation cost. If the soot aggregation was considered [282, 396, 402-404], additional transport equation for the primary particle number density is required. In addition, additional assumptions are necessary. Frequently, it is assumed that all aggregates are comprised of equal-sized spherical primary particles, and they also have the same fractal dimension $D_f$ of 1.8, while smaller primary particles are assumed to be dense spheres with $D_f$ of 3.0.
Kinetics describing the heterogeneous soot surface processes (i.e. surface growth, oxidation and condensation) are similar for both the method of moments and the sectional method, although the specific implementation is different. As mentioned previously, the soot surface growth model is mostly based on the HACA soot surface reaction scheme developed by Frenklach and coworkers [198, 280]. The kinetics of the surface reactions are described using the concept of soot surface site (C_{soot-H}). The modeling of soot oxidation by OH is based on the kinetic theory and a widely-used value of this collision efficiency is 0.13. Soot oxidation rate by O_2 can be calculated based on the concept of soot surface site, similar to the modeling of HACA kinetics. PAH addition rate can be modelled based on the collision rate between large aromatics molecules (such as pyrene) and soot particles. Note that unlike chemical growth reactions, which require surface sites, PAH addition can be regarded as a physical process such that the rates are independent on the concentration of surface species. As not all collisions lead to condensation, a collision efficiency was generally implemented to represent the probability of a successful addition of the PAH onto the soot particle surface.

We finally pointed out the importance of more recently-proposed sterically-resolved particle surface reaction modelling, as discussed in Section 2.3.

3. Experimental techniques for soot related measurements

Experimental determination for the concentrations of soot related gas- and condensed-phase species are of central importance in soot mechanism studies. Experimentation not only provides the data to validate soot models but is also a source for new insight into soot formation. As a comprehensive understanding of soot-related diagnostic methods—including their advantages and limitations—is necessary for full appreciation of the experimental data, we provide in this section a brief overview of the most widely-used experimental techniques.
3.1 Measurements of gas-phase species

3.1.1 Gas chromatography

Gas chromatography (GC) is a major technique for separating individual gaseous compounds from their complex mixtures. The most important part of a gas chromatograph is a specially coated long column through which a mixture is separated. Inside this column, gaseous compounds (mobile phase) interact with the inner coating of the column wall (stationary phase). These interactions are dependent on the physical properties of individual gaseous components; therefore, different compounds elute from the column after different retention times, leading to mixture separation. Additional sensors are required to detect and quantify the eluate, and those widely used for combustion products are the flame ionization detector (FID) for hydrocarbons, and thermal conductivity detector (TCD) for H₂, CO, CO₂, H₂O, N₂ and other non-combustible gases. In many cases, a mass spectrometer (MS) is also used as a highly effective detector, comprising a GC-MS system.

Figure 6 shows an example of a GC setup used by Gomez and coworkers [80] for the speciation of gas samples from counterflow flames. As can be seen, this particular setup adopts three parallel columns connected to different detectors. The TCD was used to quantify N₂ and O₂, which are present as large fractions of the sample. The FID was employed for the quantification of hydrocarbons as well as CO and CO₂, with the help of an upstream methanizer. The identification of the species was achieved by checking the molecule-specific mass spectrum.
A GC analysis requires probe sampling; therefore, it is an intrusive measurement. Probe disturbance is inevitable and needs to be considered, especially for small laboratory flames. The nozzle inner diameter of a typical microprobe used for product extractions is on the order of 100 μm [80, 242, 405, 406]. Although this size seems small, the presence of the probe can still cause the species spatial profile to shift up to several millimeters. In addition, special care must be given when measuring soot-laden flames since the small nozzle can potentially become clogged by particles. Another major drawback of GC analysis is that it may take up to several hours for all the species of interests to elute from the column, especially for large molecular species. This means that the GC method is an inherently off-line technique so that detection of short-lived radicals is impossible. As typically performed by Jahangirian et al. [80] and can be seen in Figure 6, gas samples, after being extracted from various locations of a flame, were stored in a set of sample loops to be analyzed overnight. If the species of interest include those with high boiling points (such as PAHs), the probe, the sample lines, and the storage loops must be heated to avoid condensation [407]. In such cases, it is necessary to determine whether thermal aging can cause variation in the species concentration. The

Figure 6. Schematic of GC setup for speciation of counterflow flames. Adapted from Jahangirian et al. [80] with permission of Elsevier.
condensation problem also put an additional limit on the maximum molecular weight that can be analyzed by GC.

3.1.2 Mass spectrometry

A mass spectrometer (MS) identifies different chemical species by ionizing neutral molecules and measuring the mass-to-charge ratios. In addition to use as a detector in a GC-MS system, a MS can also be used separately for combustion speciation, provided suitable sampling and ionization techniques are employed. Perhaps the most popular sample introduction method is through molecular-beam (MB) [408, 409]; its application in combustion diagnostics has been periodically reviewed by Biordi [410], Westmoreland et al. [411], Hansen et al. [172], and Qi [412]. Figure 7 shows an experimental setup used by Skeen et al. [413] for the chemical speciation of counterflow diffusion flames using molecular beam sampling. As can be seen, the apparatus mainly consists of a counterflow burner enclosed in a low-pressure chamber, a double-stage pumping flame-sampling system, and a time-of-flight mass spectrometer. An important advantage of the MBMS technique is its ability to preserve radicals in a molecular beam for further MS detection. Nevertheless, it is still noteworthy to point out that MB sampling, as an intrusive diagnostic method, is not used directly to sample the counterflow flames, but the material extracted from the flames via a quartz millimeter-sized probe.
Figure 7: Schematic of counterflow burner coupled to a time-of-flight mass spectrometer with molecular beam sampling and VUV photoionization. Reprinted from Skeen et al. [413] with permission of Elsevier.

After the formation of a molecular beam, neutral atoms or molecules must be ionized before they can be sorted and identified based on their mass-to-charge ratios. Various ionization methods have been developed, among which electron ionization (EI), laser-based photoionization, and synchrotron VUV photoionization are the most widely-used in combustion research. EI is based on the bombardment of high energy electrons to molecules in producing ions. EI cross-sections are a strong function of electron energy, with rather low values near the ionization threshold for most species. Therefore, for strong signals the electron energy is frequently set at a level much higher than the typical ionization energy threshold for combustion species (around 70 eV, as the maximum ionization cross-sections occur at this energy level). Unfortunately, these energetic electrons can cause significant fragmentation of the parent molecules. If the sample is a single compound, the fragmentation pattern produced provides a signature of its molecular structure and thus is helpful for the structural determination of unknown species. However, if the sample is a complex mixture, the
fragmentation may cause significant difficulties in deconvoluting individual species, if not making the process completely impossible. In this regard, MS with EI is thought to be more suitable for use as a detector for GC-MS systems where the mixtures are separated before entering MS.

To minimize or avoid fragmentation, ionizing sources with wide tunability (in the range of ionization energy of combustion relevant species, typically 7–16 eV) and high energy resolution are required. Synchrotron radiation in the VUV regime has been proven to be an ideal ionization source [412]. The photon energy can easily be tuned to just above the threshold ionization energy of most combustion intermediates so that the neutral molecules/atoms/radicals can be ionized (single-photon ionization) without significant fragmentation, resulting in much cleaner mass spectra for easy analysis. Furthermore, by recording the mass spectra of a certain species at different photon energies, a PIE spectrum can be obtained. The information about ionization energy contained in a PIE spectrum allows unknown species to be identified by comparison against values in the literature or obtained from quantum computation. Since most isomers have different ionization energy, a PIE provides an effective way to distinguish isomers. Prominent examples are the separation of fulvene and benzene (C₆H₆), allene and propyne (C₃H₄) [171], and ethenol and acetaldehyde (C₂H₄O) [414], among others.

With all its advantages, synchrotron radiation is only accessible in a limited number of locations (mostly large national laboratories) across the globe. Consequently, beam time dedicated to combustion speciation is extremely limited. As an alternative, VUV lasers can also be used for single-photon photoionization, although these laser-based methods suffer from two major drawbacks: 1) The photon energy is either fixed or inconvenient to tune and 2) Typical maximum photon energy is not enough to ionize species with high ionization energies. Bermudez and Pfefferle [415] frequency tripled the third harmonic of a Nd-YAG laser in a xenon cell to produce a VUV beam at 118 nm (i.e., photon energy of 10.5 eV), which was subsequently used to ionize most C₃ and larger hydrocarbon species (except C₃H₈ and n-C₄H₁₀) for MS analysis [416]. However, major combustion products, and most C₂ species, have ionization energy above 10.5 eV and thus cannot be photoionized. To achieve
tunability, a four wave sum-difference frequency generation in a krypton cell can be used, which typically requires two dye lasers and has been shown to produce broadly tunable VUV radiation with photon energy between 6.2 and 10.3 eV [417]. This technique has been successfully used by Qi and McIlroy [418] and Werner and Cool [419, 420] for combustion speciation.

In addition to single-photon ionization, resonance-enhanced multi-photon ionization (REMPI) has also been used in MBMS measurements [421-423]. REMPI involves the process of resonant multi-photon (typically 1 or 2) absorption by a molecule or atom to reach an intermediate Rydberg state, followed by additional photon absorption that leads to ionization. The signature vibronic transitions of individual species, together with the photon energy tunability of dye lasers, provides species selectivity. Even in cases where there is an absorption energy band overlap so that more than one species can be ionized with the same laser wavelength, the produced ion mass may be different; thus the combination of MS and REMPI offers the possibility for two-dimensional analysis. When the excitation photons and the ionizing photon are of different energies, two-color REMPI provides an additional method if the conventional, one-color REMPI technique is insufficient for species selectivity. In fact, REMPI-MS is both mass-selective and isomer-selective, due to the fact that isomers with different molecular structures have different spectroscopic features. It is believed to be especially suited for selectively ionizing and measuring aromatic species [424-428]. It is, however, worthwhile to mention that, ideally, the samples should be supersonically jet-cooled to an ultra-low rotational temperature (i.e., tens of Kelvins) before being photon-ionized, in order to reduce the broadening of energy levels which are detrimental to REMPI energy selectivity. Fortunately, this can be achieved by molecular beam sampling, although care must be taken when sampling high temperature combustion products at low pressures.

A notable advantage of REMPI, as compared to tunable laser-based VUV, is that the beam can be more easily obtained via a single dye-laser in the comfortable UV regime. The drawbacks of REMPI-MS are: 1) Many necessary spectroscopic data (i.e., energy bands, excitation cross-sections,
ionization cross-sections, and etc.) are not readily available [123]; 2) Significant experimental time is required as the photon energy (i.e., the laser wavelength) must be changed [123], and related calibration must be performed (ideally) for each species of interest; and 3) Its application is typically limited to small and medium-sized aromatic species [412]. To summarize, of the above three different ionization methods, EI is universal but not selective, REMPI is selective but not universal, and tunable VUV is both universal and selective. It is important to mention, however, that REMPI combined with MBMS is probably the method of choice [127] for quantitative PAH measurements, due to the sensitivity enhancement from resonance, which is especially important for species with ultra-low concentrations such as PAHs.

3.1.3 Optically-based in-situ methods

Both GC and MS techniques are intrusive techniques, thus they share the disadvantages of flame perturbations. The presence of the probe may affect the temperature/concentrations profile as well as the flow field [429]. On the other hand, laser-based optical methods can provide an in-situ determination of species concentrations from unperturbed (in most cases) flames.

Species-selective laser-based combustion diagnostic techniques primarily include laser absorption spectroscopy (LAS) [430, 431], laser-induced fluorescence (LIF) [432], and Raman scattering. Excellent books and reviews for these methods are available in the literature [430, 431, 433]; here only a brief introduction is offered.

A molecule with dipole moment can absorb photons that have energy resonant with the spacing between two of its different ro-vibrational states, which are the signatures of individual species (thus species-selective). The extent of absorption is proportional to the number concentrations of the absorbing species, through the Beer-Lambert law, \( I = I_0 \times \exp(-\eta \lambda L) \), where \( I \) and \( I_0 \) are the transmitted and incident beam intensities, respectively; \( \eta \) is the spectral absorption coefficients and, \( L \) is the path length. For a spectrally-isolated transition, \( \eta \) can be written as:
\[
\eta_\lambda = s(T) \times \varphi(\lambda) \times P_i
\]

where \(s(T)\) is the temperature-dependent transition line strength, \(\varphi(\lambda)\) is the line shape function depending on thermodynamic conditions, and \(P_i\) is the partial pressure (thus mole fraction) of the probed species. Detailed expressions for \(s(T)\) and \(\varphi(\lambda)\) can be found in Ref. [434]. For large molecular species that do not present a spectrally-resolved rotational transition, the overall absorption cross-section (\(\sigma_{\text{abs}}\)) can be used:

\[
\eta_\lambda = \sigma_{\text{abs}} \times N_i
\]

where \(N_i\) is the molecule number concentration. Although Eq. 11 appears to be simpler, \(\sigma_{\text{abs}}\) must be experimentally calibrated for a specific gas under specific thermodynamic conditions, and an extrapolative extension to experimentally uncovered conditions is very difficult [430].

LAS is mostly conducted in the infrared spectrum since ro-vibrational transitions typically involve energy differences corresponding to photons in the near-mid infrared regime. Since each species has specific signature transitions, tunable diode laser source is generally the workhorse for LAS, providing wavelength tunability to be resonant with such transitions, hence the term tunable diode laser absorption spectroscopy (TDLAS). LAS is especially suitable for quantitative determination of species mole fractions in 0D shock tubes, 1D laminar flat premixed flames and axisymmetric 2D coflow and counterflow flames. Under such conditions, the local absorption coefficients (\(\eta_\lambda\)) along the line-of-sight are either uniform or can be obtained through simple tomographic inversion. For unsteady multi-dimensional conditions which are predominant in practical combustion devices, the line-of-sight feature of LAS becomes a major limitation for its application.

In combustion environments where many intermediates coexist, the species to be quantified typically needs to have signature transitions (at high temperatures) that are interference-free from other species. Large molecules tend to have very complex and broad absorption spectra and will therefore almost always interfere with other species. As a result, quantifiable species quantification with LAS is most frequently done for small molecules or radicals such as \(\text{CO, CO}_2, \text{H}_2\text{O, CH}_3, \text{CH}_4,\)
C₂H₂, and C₂H₄. Nevertheless, if interference can be suitably taken care of or if qualitative monitoring is targeted, measurements using LAS can be extended to practical combustion conditions for larger molecules such as n-heptane, iso-octane, and methyl-cyclohexane [435].

Another concern for TDLAS is the lower limit for detection. If the absorbance level is too low (either due to the low concentration of probed species or its low line strength), light absorption could be too weak, which may adversely affect the accuracy of measurements. In this regard, technical approaches are available to increase the path length, or cavity ringdown spectroscopy (CRDS) can be applied, which is based on the idea to reflect the laser beam back and forth inside the cavity bounded by two highly reflecting mirrors [436]. Typical mid-IR laser sources are tunable only in a relatively narrow wavelength range. Therefore, multiple lasers are required if more than one species specific need to be measured. We quote recent studies using TDLAS to measure C₂H₂ concentration formed during the pyrolysis of ethylene and benzene in a shock tube [39], and to measure temperature via CO₂ absorption in sooting ethylene coflow diffusion flames [437] to demonstrate the potential of this method in quantifying important soot precursors as well as temperature under sooting conditions.

Laser-induced fluorescence (LIF) is a process in which the target molecule/radical is excited to a higher energy state after absorbing one or more (typically two) photons, and subsequently returned to the ground state by spontaneous emissions of photons (i.e., fluorescence). Qualitatively, species selectivity can be obtained by tuning the laser wavelength to be resonant with the target species. Quantitatively, the fluorescence intensity is proportional to the number concentrations of the probing species. A notable advantage of LIF as compared to LAS is spatial resolution: LIF is a point-based (as opposed to line-of-sight based) measurement with signals typically detected perpendicular to the incident laser beam. In addition, LIF can be configured to measure species concentration distribution in a 2D plane (planar LIF), by manipulating the excitation laser beam to form a light sheet and using an intensified charge-coupled device (ICCD) camera as a fluorescence signal detector.
Unlike LAS, LIF typically involves electronic transitions of target species so that lasers in the UV-Vis spectrum are used as sources of excitation (electronic state transitions have energy differences corresponding to photons in the UV-Vis spectrum). This is due partly to the relatively large Einstein A coefficients of electronic transitions and partly to the availability of high-powered UV and visible lasers. In combustion studies, OH, CH, H radicals and NO and CO molecules are frequently measured with LIF. Note, however, due to the uncertainties in fluorescence quenching processes and the fluctuations of laser energies, quantitative determination can be challenging even for these small species. On the other hand, LIF is especially suitable for qualitatively tracking the spatial distribution and temporal evolution of flame front (as marked by CH or OH). By seeding fluorescent tracers (i.e., acetone), LIF is also frequently used to visualize the mixing process of two streams.

In addition, LIF is used to measure PAH concentration in a qualitative manner. PAHs have somewhat complicated molecular structures and thus have fluorescence with broad and complex spectrum (further complicated because both temperature and surrounding atmosphere would considerably alter the spectra). In flame environments, the detected signals always represent the fluorescence of a mixture of PAH species, making quantitative deconvolution highly challenging, if not entirely impossible. Nevertheless, it has been shown that larger PAHs are better represented by LIF signals detected at longer wavelengths [438-440] so that a spectrally-resolved LIF signal can be used to qualitatively differentiate between small and large PAHs. The signal intensity can also be used to qualitatively evaluate the concentrations of these different classes of PAHs. This method is particularly useful in evaluating fuel effects on PAH formation by comparing the relative concentrations of PAHs as represented by LIF signal intensity among flames with different fuels or fuel additives [441-447]. Nevertheless, it is important to note that PAH LIF signal can be highly dependent on temperature [448]. As a result, quantitative comparisons of PAH LIF signals can be performed only when the temperature is the same, and care should be taken when evaluating LIF signals measured in areas with significantly different local temperatures. Furthermore, as mentioned
above, many different PAH species can coexist in flame conditions. The fluorescence spectra of these PAH species are not only complex and broad, they can also significantly overlap with each other. Collisional quenching with surrounding gases serves as additional source of complications for quantitative PAH LIF analysis [449].

Nevertheless, there is recent progress towards *quantitative* measurement of PAH, using LIF. The new technique—called *jet-cooled* LIF—is based on the extraction of flame-generated species through a microprobe and subsequent expansion via a supersonic jet, before being analyzed using LIF in an analysis chamber [55, 450, 451]. The supersonic expansion reduced the temperature and also the possibility of an otherwise large number of vibronic transitions, making the fluorescence spectra more structured and amenable to selective detection. Furthermore, the LIF quenching effect can also be reduced, making quantification easier. Although this method requires probe sampling and thus is not non-intrusive, it provides an important alternative for quantitative PAH measurement in flames. Nevertheless, to date, the jet-cooled LIF has been successfully used only to measure a couple of PAH species and further efforts are required to extend its range of application and to quantify measurement uncertainties.

LIF can also be used to detect flame-generated nanoparticles which exhibit molecular characteristics [51, 290, 452, 453]. Differentiation between particles and PAH molecules can be obtained by time-resolved fluorescence polarization anisotropy [454-456].

More recently, infrared LIF (IR-LIF) has attracted interests since it can be used to measure important molecules such as CO₂, CO and H₂O, which are not accessible by conventional UV-Vis LIF techniques [457, 458]. The application of this technique in harsh combustion environments is expected to be promising, especially considering that it can also provide 2-D temperature distribution using a two-color method [457] (conventional two-color OH PLIF for temperature measurements is restricted to regions where OH are abundant). We finally quote a recent study using femtosecond
two-photon laser-induced fluorescence (fs-TPLIF) to measure H atom in sooting premixed flames [459] to demonstrate the applicability of LIF in sooting flames.

Another species-selective technique frequently employed in combustion research is Raman scattering, which is based on inelastic interactions (as opposed to the elastic Rayleigh scattering) between incident photons and target molecules. The scattered light has a species-dependent fingerprint frequency shift relative to the incident light (Raman shift), which can be used to identify and quantify species. Spontaneous Raman scattering is suitable for point measurements of major combustion species such as O₂, CO, CO₂ and H₂O. An important advantage of spontaneous Raman scattering is its ability to measure multiple species simultaneously [460-462]; while an important disadvantage is its very weak signal. In practice, high-energy pulsed laser beam is focused into a measurement point and the resulting Raman scattering signal is spectrally-resolved with a spectrograph before being recorded with an intensified camera. One major drawback of Raman scattering is the costly and bulky instruments (high-resolution spectrometer, high-energy laser and intensified detectors) it requires, which limit its application largely to laboratory conditions. Note, the Raman scattering signal intensity can be enhanced by tuning the incident photon energy resonant with a specific species of interests (i.e., resonance Raman scattering) [463].

Both the LIF and Raman scattering techniques discussed above are incoherent methods and the signals leave the sample in all directions, covering the whole 4π solid angle, while the collection of the scattered signal can only be achieved over a relatively small solid angle. This reduced signal collection efficiency can be overcome by using coherent techniques in which the signal becomes beam-like and can be effectively directed to a detector. A thorough discussion of four-wave mixing techniques, among which the widely-used coherent anti-stokes Raman spectroscopy (CARS) is a particular configuration, was provided by Kiefer and Ewart [433]. Recent studies using CARS to measure temperature in sooting coflow diffusion flames [464, 465] are quoted here to demonstrate its applicability in sooting environments.
3.2 Soot measurements

Soot loading in a flame is typically represented by soot volume fraction (SVF, or \( f_v \)), so its experimental determination is central to soot research. Therefore, it is worthwhile to provide a discussion here on various diagnostic techniques developed for quantitative SVF measurements.

3.2.1 Light extinction

Soot particles are known to absorb light in a broad spectral range from UV to IR, so that the intensity of any light beam (visible or near IR) passing through a soot-laden flame is expected to be attenuated. The extent of such intensity reduction can be correlated to the volumetric loading of soot particles. In typical light extinction experiments, the intensity of a probe beam with a fixed wavelength (\( \lambda \)) is measured both before (\( I_{\lambda,0} \)) and after (\( I_{\lambda} \)) passing the flame, and their ratio is defined as the light transmissivity (\( \tau_{\lambda} \)). The local extinction coefficient \( K_{\text{ext},\lambda} \) can then be obtained through the Beer-Lambert law as:

\[
\tau_{\lambda} = \frac{I}{I_0} = \exp \left( -\int_0^L K_{\text{ext},\lambda} ds \right)
\]

Eq. 12

where \( L \) is the length of light path. For a homogeneous system in which \( K_{\text{ext},\lambda} \) is a constant along the optical path, Eq. 12 can be reduced to a simpler form:

\[
K_{\text{ext},\lambda} = -\ln(\tau_{\lambda}) / L = -\ln(I/I_0) / L
\]

Eq. 13

For axisymmetric cases such as counterflow or coflow diffusion flames, an inversion algorithm [466, 467] can be used to derive local \( K_{\text{ext},\lambda} \) from multiple realizations of line-of-sight measurements.

The inference of SVF from measured local extinction coefficients requires fundamental knowledge of soot-light interactions. The complex soot morphology makes it computationally demanding to use exact theories to analyze such interactions. Consequently, researchers frequently use the Rayleigh-Debye-Gans–fractal aggregate (RDG-FA) theory as an approximation of the otherwise intensive analysis. According to the RDG-FA theory, the local extinction coefficient can be written as:
\[ K_{\text{ext},\lambda} = N_{pp} \pi^2 D_{pp}^3 E(\tilde{m}_{\lambda})/\lambda \]  
\text{Eq. 14}

where \( N_{pp} \) and \( D_{pp} \) are the number density and the diameter of primary particles, respectively. \( \tilde{m}_{\lambda} \) is the wavelength (\( \lambda \))-dependent soot complex refractive index and \( E(\tilde{m}_{\lambda}) \) is the light absorption function, which is defined as:

\[ E(\tilde{m}_{\lambda}) = -\text{Im}\left(\frac{\tilde{m}_{\lambda}^2 - 1}{\tilde{m}_{\lambda}^2 + 2}\right) \]  
\text{Eq. 15}

Since in the RDG-FA theory, soot particles are treated as fractal aggregates of mono-dispersed spherical particles in point contact, \text{Eq. 14} further shows the total volume of the soot aggregates (i.e., soot volume fraction, \( f_v \)) to be:

\[ f_v = \left(\frac{\lambda K_{\text{ext},\lambda}}{6\pi E(\tilde{m}_{\lambda})}\right) \]  
\text{Eq. 16}

Note that in writing \text{Eq. 14}, it was assumed that light extinction is solely due to soot absorption. In reality, however, both particle scattering and light absorption by gas-phase species (such as PAHs) may contribute to intensity attenuation of the transmitted light. The ratio of scattering coefficient to absorption coefficient can be determined, provided the morphology of particles is known (see, e.g. Ref. [468]). Nevertheless, when the optical sizes of soot aggregates are significantly smaller than the laser wavelength \( (2\pi R_g/\lambda < 0.3 \), where \( R_g \) is the radius of gyration of agglomerate), light scattering becomes negligible compared to absorption [469, 470]. The absorption by a gas-phase species can be estimated by running the HITRAN code for typical combustion products. Simonsson \textit{et al}. [471] found that for a rich-premixed flame, molecular absorption in the near-infrared regime (700–1064nm, typically used for soot measurement) from major combustion products, is dominated by H\textsubscript{2}O molecules at 978.6nm. However, the absorption strength was rather small and its contribution to soot light extinction was concluded to be marginal [471]. This is especially true in practical cases where it is rare for the light wavelength used for soot measurement to be coincident with the narrow spectral line resonant for molecular absorption.
On the other hand, PAHs, with their complex molecular structures, are known to absorb significantly in the UV-Vis spectral range [472]. Since PAHs are precursors to soot, flames with high soot loadings are also featured with high concentrations of PAHs, so that PAH interference is expected to become worse for more sooting flames. If the wavelength of the probe light lies in such a UV-Vis spectrum, the interference of soot extinction from PAH absorption must be accounted for. The simplest, and perhaps the most effective remedy, is to perform soot extinction measurements with light wavelength in the near-infrared regime (i.e., >700 nm), for which PAH absorption is minimal [473]. The employment of a near-infrared beam may have additional benefits as the absorption ratio for soot aggregates is predicted to decrease with increasing λ [127]. However, it is worthwhile to point out that previously many studies in the literature used light in the visible spectrum for soot extinction measurement. Thus, this fact should be considered when interpreting data in the literature.

A major source of uncertainty in light extinction measurement is the quantitative variations of the soot complex refractive index $\tilde{m}_d$ (and thus $E(\tilde{m}_d)$) on wavelength, and on the physical/chemical properties of soot produced from different sources. The spectral dependence of $E(\tilde{m}_d)$ for mature soot from coflow ethylene diffusion flames has been studied using laser-induced incandescence (LII, discussed later) with both 532 and 1064 nm beams. It was determined that the absorption cross-sections have a $\lambda^{-0.83}$ dependence [474]. Cleon et al. [475] determined $E(\tilde{m}_d)$ along a rich premixed burner-stabilized methane flame and found that the ratio of $E(\tilde{m}_{532nm})/E(\tilde{m}_{1064nm})$ decreased significantly with the height above the burner (HAB) in the nascent soot region, but it tended to level off at higher HAB. This indicates that soot at different flame locations have different optical properties. Nevertheless, the optical properties of mature soot appear to have a more consistent value [127].

In many cases, a narrow laser beam is sent through a soot-laden flame and a single-point photodiode is used to detect the transmitted beam intensity. For a one-dimensional soot field (in a
burner-stabilized flat premixed flame, or planar counterflow diffusion flames), the spatial resolution of measurement is then determined primarily by the beam size. The spatial variation of the soot field is obtained by passing the beam through different locations of the flame, realized by either moving the flame relative to a fixed beam [140, 476], or moving the complete optical setup relative to a fixed flame burner [69]. A major disadvantage of such line-by-line measurements is that it may take a longer time to scan the entire flame, thus it is not practical to measure soot fields that are time dependent. Alternatively, a 2D collimated beam that covers the entire flame area (projection along the direction of the beam) can be employed, with a camera serving as the device for light intensity detection [477, 478]. In such a configuration, every pixel of the camera records the intensity of a light ray transmitted through a different location of the flame. Hence, a complete map can be obtained at once, allowing for instant measurements of time-varying soot fields, for example, in flickering coflow diffusion flame [479]. Note that beam steering, which is caused by the lensing effect of flame gases, must be minimal or its influence should be accounted for in extinction measurements, especially in cases where a large temperature gradient (therefore a large density gradient) exists. Examples of such cases include counterflow diffusion flames and high-pressure coflow flames [477].

3.2.2 Spectral soot emission

The spectral emissions of flame-heated soot particles provide information on both soot temperature and volume fractions. In axisymmetric flames, soot radiation transmitted along a line-of-sight can be detected and spectrally resolved using a spectrometer. A typical optical setup for such spectral soot emission (SSE) measurement is schematically shown in Figure 8. The detected light intensity can be described as [480, 481]:

$$I_\lambda(y_0) = \int_{-\infty}^{\infty} \eta_\lambda(x, y_0) B_\lambda[T(x, y_0)] \exp(-\int_0^{\infty} K_{\text{ext},\lambda}(x', y_0)dx') \, dx$$  \hspace{1cm} \text{Eq. 17}$$

where $y_0$ denotes the distance between the line-of-sight chord and the symmetry plane of a flame/burner; $x$ is the distance along the chord under consideration, with 0 defined at the symmetry
point; $\eta_\lambda$ is the local soot absorption coefficient, and is equal to the soot extinction coefficient $K_{\text{ext},\lambda}$ under the assumption of negligible scattering [482], so that it is related to SVF through Eq. 16. $B_\lambda$ is the Planck function of wavelength $\lambda$ and temperature $T$ and is expressed as:

$$B_\lambda(T) = \frac{2\pi c^2}{\hbar^5} \frac{1}{\exp(hc/\lambda kT) - 1} \approx \frac{2\pi c^2}{\hbar^5} \exp \left(-\frac{hc}{\lambda kT}\right)$$

Eq. 18

where $h$, $c$, and $k$ are the Planck constant, light speed, and Boltzmann constant, respectively. Note that the Wien approximation to the Planck’s law (rather accurate in the visible to near infrared spectrum) is invoked in writing in the latter half of Eq. 18. In Eq. 17, the term of $\eta_\lambda B_\lambda$ (i.e., $K_{\text{ext},\lambda} B_\lambda$) represents the local spectral emission, and the exponential term takes the form of the Beer-Lambert law to describe the attenuation of such emission by self-absorption along the line-of-sight.

Figure 8. Typical optical setup for spectral emission–based soot temperature and volume fraction measurements (a), reprinted from Thomson et al. [483] with permission of Elsevier and schematics.
of line-of-sight emission intensities ($I_b$) for axisymmetric flames (b), reprinted from Liu et al. [484] with permission of Elsevier.

Through horizontal translation of the flame relative to the detection system, line-of-sight measurements can be performed at different $y_0$, and a subsequent tomographic inversion will provide local values of $K_{ext,\lambda}(r)B_\lambda(r)$ [484]. Substituting Eq. 14 into Eq. 18, taking the logarithm on both sides, and considering $K_{ext,\lambda}(r)B_\lambda(r)$ is now known at all radial positions $r$ lead to [484]:

$$\ln \left[ \frac{E(\tilde{m}_\lambda)}{K_{ext,\lambda}(r)B_\lambda(r)\lambda^2} \right] = \frac{hc}{\lambda kT(r)} - \ln[12f_v(r)\pi^2 hc^2]$$  \hspace{1cm} \text{Eq. 19}

As seen, the left side of Eq. 19 has a linear relationship with $1/\lambda$ and the slope, $hc/kT$, is solely determined by temperature. In practical applications, $hc/kT$ can be obtained by a curve fitting using the method of least mean squares. Once the temperature is known, Eq. 19 can be used again to provide $f_v$.

One practical challenge in this spectral soot emission technique is the accurate determination of the local values of $\eta_\lambda$, or equivalently $K_{ext,\lambda}$, with the assumption of zero scattering. First, the detection system must have a large depth of field to allow every emitter along the line-of-sight to be properly focused into the spectrometer. This was shown to be achievable using an imaging lens with a sufficiently large f-number [480] (for example, f/48 was used in recent studies [485, 486]). Second, the self-absorption of soot emission must be accounted for [480] because: 1) Emissions from different emitters along the line-of-sight will be attenuated unequally due to the different path lengths toward the detector, causing the axisymmetric assumption needed for Abel-like tomographic inversion to break down and 2) Self-absorption is also dependent on $\lambda$ and as such it complicates the soot temperature measurement by shifting the slope, as discussed in Eq. 19. In view of these issues, Snelling et al. [480] proposed an iterative method to correct for self-absorption, and Liu et al. [484] concluded that when the maximum flame optical thickness exceeds 0.3 such correction becomes particularly important.
Also suggested by Eq. 19, the emissions theoretically need to be detected at only two wavelengths to find the flame temperature and $f_v$, considering two points in the $(\frac{E(m)}{K_{ext}(r)B(\lambda r)\lambda^4}, 1/\lambda)$ plane can already determine the line whose slope is $hc/kT$. The experimental technique based on this fact is known as two-color pyrometry and it was also used extensively in soot measurements. Although the measurement uncertainty can be larger than its multi-wavelength counterpart, a spectrometer is no longer needed and 2D measurements are possible with a color filtered CCD camera as a detector [487, 488].

This spectral soot emission (SSE) technique has been applied successfully in many soot studies, especially for coflow diffusion flames at high pressures (see Karatas and Gülder [107] and references therein). A notable advantage of SSE is that no additional illuminating source is required, and for high-pressure applications, optical access is needed only from one side of a high-pressure chamber, simplifying the design of the pressure chamber [107]. It is noted, however, that SSE was employed for soot studies in counterflow diffusion flames only very recently by Gleason et al. [489, 490]. By contrasting the soot temperature as determined by SSE against the well-established temperature from the literature, the authors were also able to obtain information on the dispersion exponent, a soot optical property related to its hydrogen content.

3.2.3 Laser-induced incandescence

Laser-induced incandescence (LII) refers to the quasi-blackbody radiation of soot particles after being heated up to about 4000 K by high-energy laser beams. Through analysis of such radiation, many properties of soot ensemble can be derived, such as SVF and soot primary particle size. Dedicated reviews on the LII technique for soot diagnostic exist in the literature [126-128, 491], and a brief overview of the basics of this promising method is given here.
The usefulness of LII stems from the fact that SVF is reasonably proportional to LII signal intensity \((I_{LII})\) [492, 493]. Melton [492] has shown that, for soot ensembles with primary particle sizes smaller than the incident laser wavelength, the LII signal intensity can be written approximately as:

\[
I_{LII} = C_1 \int_0^\infty N P(D) D^n dD, \quad \text{with } n = 3 + 154/\lambda_{det}
\]

Eq. 20

where \(N\) is the total particle number density, \(P(D)\) is the normalized probability density for particle with diameter \(D\), and \(\lambda_{det}\) is the detection wavelength in nm unit. As can be seen, for larger detection wavelengths, the exponent \(n\) approaches three and \(I_{LII}\) becomes linearly proportional to SVF.

It is noteworthy that Eq. 20 was obtained in the limit of high laser power. Whether this limit is achieved in practice can be tested by plotting \(I_{LII}\) as a function of laser pulse energy. Typically, \(I_{LII}\) initially increases, reaching a plateau regime (saturation), then begins to decrease again (due to soot evaporation) with progressive increases in laser energy [494]. Experiments are typically conducted in the plateau regime so that the signals are not affected by the pulse-to-pulse energy variations of the laser beam. Due to potential interference from PAH fluorescence, it is generally recommended that infrared light be used as the excitation source [472], although a recent work [459] showed that by delayed detection, the SVF measured using UV (266 nm) excited LII is similar to that excited by a near-infrared beam (1064 nm).

LII measurement alone cannot provide the absolute value of SVF, since the proportionality constant \((C_1\) in Eq. 20\) depends on many parameters requiring experimental calibration. Additional light extinction/spectral emission measurements [472], or absolute irradiance calibration [495, 496], are possible approaches that provide quantitative relations between LII intensity and absolute SVF.

When the LII signal is analyzed in a time-resolved manner, information about particle size can be obtained in addition to SVF. This is based on the fact that particles with different diameters are cooled down (after laser heating) at different rates [497]. Large particles have low surface area to volume ratio and thus are cooled relatively slowly compared to small particles. To obtain the quantitative relationship between LII signal decay rate and soot particle size, a model that describes the heat/mass
exchange processes between the particle and the surrounding gases [498] is required. There are three major paths for energy loss: sublimation of soot particles, heat conduction to the surrounding gas and radiation of soot particles. Such energy losses can be corrected with the internal energy of soot particles through the following energy conservation equation [126]:

\[
\frac{dU}{dt} = \dot{Q}_{abs} + \dot{Q}_{cond} + \dot{Q}_{sub} + \dot{Q}_{rad}
\]

Eq. 21

where \(U\) is the internal energy of soot particle, \(\dot{Q}_{abs}\) is the rate of laser energy absorption, \(\dot{Q}_{cond}, \dot{Q}_{sub}\), and \(\dot{Q}_{rad}\) are the rates of energy loss due to heat conduction to surrounding gas, sublimation and heat radiation, respectively. The rate of change of soot internal energy can be written as:

\[
\frac{dU}{dt} = \frac{\pi}{6} d^3 \rho_s c_s \frac{dT_s}{dt} = m_p c_s \frac{dT_s}{dt}
\]

Eq. 22

where \(\rho_s, c_s,\) and \(T_s\) are the density, specific heat, and temperature of soot, respectively, and \(m_p\) is the mass of primary particle. The detailed size-dependent expression for \(\dot{Q}_{abs}, \dot{Q}_{cond}, \dot{Q}_{sub}\) and \(\dot{Q}_{rad}\) can be found in the review by Michelson [126], and they are generally dependent on primary particle sizes.

An important parameter in determining heat loss rates is soot temperature, which can be obtained by taking the ratio of soot radiation intensity at two different wavelengths (i.e., two-color pyrometry) [499]. With all the above information, a theoretical decay rate of soot radiation intensity can be computed, and comparison with an experimentally obtained rate would provide information on the primary particle size \((D_{pp})\) of soot under investigation.

In practice, time-resolved LII signals can be recorded by a photomultiplier tube (PMT) with low rise-time to provide high spatial resolution. Multiple PMTs can be used with different spectral filters installed to detect LII signals at different wavelengths, providing spectral information necessary for the retrieval of soot temperature [500]. Alternatively, planar LII can be performed using a high energy laser sheet as the excitation source and an ICCD camera as a signal detector. In this case, temporal resolution can be achieved by making measurements with consecutive gate delays in steady flames [501] and a single-shot measurement is also possible by using multiple ICCDs preset with different
delay time [502]. The latter technique is essential for measurement in highly-transient turbulent flames [503].

3.2.4 Thermophoretic sampling

In addition to the optical techniques discussed above, thermophoretic sampling is another important method for obtaining information regarding size distributions, nanostructures, chemical compositions and various other important properties of collected soot particles. The thermophoretic effect refers to the drift of particles in a direction from high- to low- temperature regions. Thermophoretic sampling makes use of this particle drift to collect hot soot particles on cool probes. The collected particles can then be analyzed by various methods, including TEM and AFM. The rate of deposition can also be linked to local SVF so that accurate determination of the total mass of the sample can be an effective method for SVF measurement, as demonstrated by Köylü et al. [504]. This method is particularly attractive as it can determine SVF without prior knowledge of soot optical properties such as a refractive index.

One important consideration when performing thermophoretic sampling is the dwelling time of the probe inside a flame. If the dwelling time is too short, not enough particles can be collected for analysis. If the probe spends too much time in the flame, processes like thermal annealing may change the properties of the collected particles and the high temperature of flame may also break some more fragile probes, such as a TEM grid. In practice, rapid insertion of a probe through double-acting pneumatic, or electric-driven piston, is frequently used and the dwelling time can be accurately controlled to the range of tens of milliseconds [485, 505]. In weakly sooting conditions, multiple insertions can be performed to collect enough soot samples [505].

It is noted that thermophoretic sampling is seldom performed in CDFs [506] compared to its wider application in coflow or premixed flames. This could be explained by the fact the very narrow sooting region (~ 1 mm) in CDF makes spatially-resolved thermophoretic sampling using conventional TEM
grid very difficult, especially considering the large vibration of a probe (to which the TEM grid is attached) after its rapid insertion (confirmed by high-speed images [507]). An innovative solution could be to insert a fine SiC wire into a flame, with a small angle between the flame axis and the wire (tilted wire), as pioneered by Figura and Gomez [508] in the PhD thesis of Figura. In this way, the wire could cover the entire axial extent of the flame, and particles in different axial positions of the flame will deposit on different parts of the wire. The disadvantage with this approach is that, due to the presence of the SiC wire (which blocks the electron beam in the TEM analysis), the sampled particles can only be viewed from the side. Recently, Carbone et al. [54] improved this SiC wire sampling technique and successfully applied it in a premixed flame, producing quantitative results (size distribution and volume fraction) via scanning electron microscopy (SEM) and helium ion microscopy (HIM) analysis of the soot samples.

Particle sampling is inherently more difficult for high-pressure flames due to the presence of a pressure chamber and associated limited physical access. The samples need also be transferred out of the chamber for further analysis after each collection, which may involve time-consuming depressurization and re-pressurization procedures. To address this issue, Vargas and Gülder [509] developed a multi-probe sampling system housed inside the pressure chamber including a circular sampling disk with ten extent-adjustable probe arms. A motor drive with programmable control unit determines which probe is used and the circular velocity of the sampling arm—and thus the sampling time. After each sampling process, the relevant probe can be retracted and the samples stored. When all the probes are loaded by samples, the experiment can be stopped, and the chamber is depressurized for sample retrieval.

A TEM analysis of sampled particles can provide information on the morphology of the soot aggregate and sizes along with the internal structures of primary particles [510]. In addition, if enough aggregates are sampled, statistical information on the particle size distributions can also be obtained. These are important information complementary to what can be obtained from optical techniques.
Nevertheless, it is worth pointing out that through a combination of multiple diagnostic methods, Zhao et al. [505] demonstrated that nascent soot particles are liquid-like and are largely transparent to the electron beam of TEM. As a result, the TEM image contrast of these nascent particles is rather small. In addition, the evaporation of soot materials, as well as carbon crystallization and annealing, was also observed to be caused by the interaction of the soot sample and the high-energy electron beam of TEM [511, 512]. These side effects make it a challenge to measure nascent soot particles with diameters below 10 nm with TEM [511]. Furthermore, as the nanostructure of nascent particles may be altered under the irradiation of high-energy electrons, results from HRTEM should be analyzed with caution.

HIM is a promising alternative to TEM in measuring nascent soot particles. With much less damage or alteration made to the sample, HIM can detect nascent soot particles with higher contrast [54, 511]. Furthermore, HIM was reported to identify particles as small as 2 nm [511], notably extending the lower size limit detectable by TEM (typically 10 nm). Nevertheless, it should be noted that HIM cannot provide information of the inner structures of samples.

If larger amounts of soot particles are required, a water-cooled suction tube can be used, and the particles deposited on the cool tube walls, or filters placed inside the tube can then be extracted by solvents [125]. The soluble material can be studied by liquid chromatography, UV absorption and fluorescence [125] while the particles can be further analyzed by—for example—a thermogravimetric analyzer (TGA) for the oxidation characteristics, X-ray diffraction (XRD) for crystalline structures, and Raman spectroscopy and Fourier transform infrared spectroscopy (FTIR) for chemical properties.

3.2.5 Differential mobility analysis

Differential mobility analyzer (DMA) is becoming an increasingly popular tool for the determination of size distributions of flame-generated soot particles [18, 49, 52, 54, 125, 355, 513-515]. DMA, as a size-selecting device, is usually adopted in combination with a particle counter,
constituting a *scanning mobility particle sizer* (SMPS) system. In application, soot-laden flame gases were first sampled and diluted within a probe and then charged by a neutralizer with known charge distributions. In an external electric field, the charged particles would move to an electrode of opposite charge at different speeds that are dependent on their electric mobility and the electric field strength. By scanning the strength of the external electric field, particles with different mobility sizes can be separated in a similar manner as in a mass spectrometer, and subsequent measurement in a particle counter provides number concentrations of the size-selected particles. After correcting the mobility size for true particle size [516-518], a size distribution can be obtained.

Either an electrometer [54] or a laser optical counter (based on light scattering by particles) can be used to determine the number density of the classified particles. In the latter case, condensation growths of these tiny particles are required before they can be counted optically, due to the insensitivity of the laser-based method to nano-sized particles. Condensation particle counter (CPC) [519] is widely used for effectively counting nano-sized particles and a detailed description of its working principle can be found in Ref. [520]. A brief description is as follows. A particle-laden flow is brought into contact with a supersaturated vapor of a working fluid, which condenses on the particles so as to increase their sizes to an extent enough for optical detection. The level of supersaturation of the condensing vapor is of critical importance to the lower detection limit of particle sizes. With the increase in the supersaturation level, progressively smaller particles can be activated. On the other hand, however, if the supersaturation level is too high, homogeneous nucleation of the vapor itself will take place, interfering with the target particles and thus leading to false counts. Iida et al. [521] pointed out that the use of working fluids with high surface tension and low saturation vapor pressure is helpful to extend the lower limit of detectable particle size. Commercial instrument typically uses water, iso-propanol or n-butanol as the working fluid with detection limit down to 2.5 nm (e.g., TSI 3776 Ultrafine CPC with n-butanol). More recently, a diethyl glycol (DEG)-based condensation nano-enhancer, when used in combination with a second-
stage growth booster, has been demonstrated to be capable of detecting particles with sizes down to 1 nm [522]. Through a theoretical investigation, de la Mora [523] anticipated that water at low temperature could even activate particles with diameter as small as 0.6 nm.

An important consideration when using an SMPS to measure probe-sampled soot particles is to avoid particle losses (e.g., through particle coagulation or diffusion to the wall) in the sampling probe and transfer tubes. For example, it was shown that a loss of 10% of particles can occur in 20 ms for a typical flame gas that contains $10^{10}$ particles per cm$^3$ [318]. As such, immediate dilution of the sampled gas is of paramount importance [49]. This means that a large amount of diluent must be mixed with the sampled flame gas immediately, posing significant challenges for the design of a sample system. A proven methodology was demonstrated by Wang and coworkers [48, 97, 230], who utilized a tubular probe, crossing a flame with a small pin hole drilled to sample the flame gas, while a carrier gas of fixed flow rate served as the diluent. Since a large flow rate of the diluent is required, the tubular probe must be relatively large (typically a tube with 1/4-inch OD is used) so that it may significantly affect the flow field and the flame. To address this issue Abid et al. [48] proposed, for applications in burner-stabilized flat premixed flames, to conceal the probe in a stagnation plate, which also serves as a stabilizing plate and a downstream boundary. As long as this downstream boundary can be accounted for in accompanying numerical simulations, the presence of the probe posed no problems. Nevertheless, this proposed method is only applicable to premixed burned-stabilized flames. For other types of flames, significant challenges still exist in the design of suitable sampling systems with minimal influence on the flame itself. This challenge may explain why the majority of SMPS flame measurements were performed for premixed flames, while limited studies for coflow diffusion were mainly performed at the flame tip [515].

Particle size distribution, measured using SMPS, can be a rich source of information on various soot formation processes, as detailed in a previous section. Singh et al. [322] (also summarized in Ref. [18]) showed that various features of a size distribution function are directly related to--for
example—nucleus size, nucleation, surface growth, and coagulation rates. In this regard, and also as pointed out by Wang [18], many soot model parameters can be fine-tuned and made more practical by validation against the experimentally measured size distributions.

There are additional techniques for soot characterization, such as dynamic light scattering, X-ray scattering, neutron scattering, aerosol mass spectrometry, etc. The scope of this review does not allow for further discussion of these techniques. More details can be found in excellent summaries by Kohse-Höinghaus and Jeffries [524], D’Anna [125], Wang [18], Desgroux et al. [127], and Michelsen [128].

4. Sooting structures and soot evolution in counterflow flames

Knowledge about sooting structures is necessary for a fundamental understanding of soot evolution in flames. As such, we discuss in this section the general flame and sooting structures in CDFs, along with their differences/similarities to those in coflow diffusion flames. We then highlight the response of soot evolution to flame structure variations, which is followed by a discussion on the effects of strain rate on sooting processes in CDFs. This section concludes with a description of soot formation in partially-premixed counterflow flames.

4.1 Sooting structures of CDFs and comparison with coflow diffusion flames

In a counterflow flame, the nozzle exit velocities for the fuel and oxidizer streams are with opposite signs. Therefore, somewhere between the nozzles there always exists a location with zero axial velocity, that is the stagnation plane. The relative positions between the high-temperature reaction sheet and the stagnation plane have a strong influence on soot evolution in CDFs. Specifically, sooting CDFs can be differentiated into two distinctive types based on this relative position [73]: soot formation flames and soot formation/oxidation (SFO) flames. The schematics shown in Figure 9 help to clarify this point.
Figure 9. Schematics of evolution for soot formation (SF) flame located on the oxidizer side of the stagnation plane (a) and soot formation/oxidation (SFO) flame on the fuel side of the stagnation plane (b). Mie scattering images are also included to show soot leakage (thus emission) through the particle stagnation plane in SF flame. In SFO flame, initially formed soot can be oxidized completely, so no soot emission can be observed [73].

Figure 9a depicts a situation when the flame is located on the oxidizer side of the stagnation plane. In such a case, soot particles, which are incepted on the fuel side from the flame sheet, will be convected away from the flame towards the stagnation plane, without passing through the oxidative high temperature flame region. While being transported, these soot particles will continue to grow through surface reactions and particle-particle coagulations. They will eventually leak out radially to the environment, because it is unlikely for these large particles to diffuse across the stagnation plane. Soot oxidation is not present during this process, since oxidizing species such as OH and O$_2$ only exist with considerable concentrations on the oxidizer side of, or near, the flame sheet. As a result, such flames are termed soot formation flames. In contrast, when the flame sheet is located on the fuel
side of the stagnation plane (Figure 9b), soot particles are convected towards the flame, where intensive soot oxidation by O\textsubscript{2} and OH radicals occur. In many cases, the initially formed soot particles can be completely oxidized, so no soot emission can be observed out of the flame. This type of CDF is termed the soot formation/oxidation flame.

More detailed flame and sooting structures for SF/SFO flames were investigated by Kang et al. [73]. The axial profiles of temperature, fuel, oxygen, and important soot precursor concentrations, as well as those of SVF, average soot particle size and number density, are shown in Figure 10 for typical SF and SFO flames fueled by propane.

For the SFO flame ($X_{f,0} = 0.25$ and $X_{O,0} = 0.7$, where $X$ is the mole fraction, the subscripts F and O indicate fuel and oxygen in the fuel and oxidizer streams, respectively; the subscript 0 is the freestream, and $z$ is the distance from fuel nozzle) shown in Figure 10(a-d), the location of the flame sheet (represented by the maximum temperature location $z_{T,max}$) stays close to the stagnation plane ($z_{stg,p}$). Here, the difference between the particle and gas-phase ($z_{stg}$) stagnation planes can be attributed to the thermophoretic effect. Fuel pyrolysis and PAH growth occur on the fuel side of the particle stagnation plane (also of the flame sheet), confirmed by the profiles of C\textsubscript{2}H\textsubscript{2} mole fraction (a) and the PAH fluorescence signal $Q_f$ (b). The sooting zone, indicated by the light scattering signal $Q_{VV}$, is also located in this region (b). It is noted that $Q_{VV}$ begins to rise near $z = 5$ mm (~1700 K), indicating the initiation of soot inception at this position. The coincidence of the location of peak PAH fluorescence (represented by $Q_f$) with the location at which soot inception begins may indicate the critical role of PAHs for soot inception. As the formed particles are convected (to the right) towards the particle stagnation plane, their number density continues to decrease (c), due primarily to particle-particle coagulation. Meanwhile, the average particle size $D_{63}$ (c) and soot volume fractions increase through surface growth and coagulation (d), reaching their peaks near $z = 6$ mm, and then decreasing due to oxidation. Note an appreciable amount of OH radicals exists on the fuel side of the flame sheet (a), where the soot zone is situated.
Figure 10. Flame and sooting structures of SFO (a,b,c,d) and SF (e,f,g,h) flames, adapted from Kang et al. [73] with permission of Elsevier.

In the SF flame ($X_{F,0} = 1.0$, $X_{O,0} = 0.25$) shown in Figure 10(e-h), the flame front is spatially separated from, and on the oxidizer side of, the stagnation plane. $C_2H_2$ and PAH fluorescence are distributed between the stagnation plane and the flame sheet (e,f). The peak of the number density profile and the onset of $Q_{VV}$ are located near $z = 7$ mm, where soot inception occurs. These newly-incepted soot particles are then convected towards the stagnation plane (away from the flame sheet). Following the direction of this convection transport (to the left), soot number density (g) decreases, but average particle size (g) and soot volume fraction (h) increase, through both coagulation and surface growth, reaching their peaks at the stagnation plane. In this process, soot oxidation hardly occurs because few OH radicals exist in the sooting zone of $z = 5–7$ mm (e). Note, it was observed [242, 281, 407, 525, 526] that a SF flame visually exhibited three distinct zones from the fuel side to
the oxidizer: an orange zone, a luminous yellow zone on the fuel side of the flame, and a blue flame reaction zone. This is consistent with the flame and soot zone structures shown in Figure 10 (e-f).

The above work of Kang et al. [73] offers general knowledge on sooting characteristics of SF and SFO flames, which were further investigated through measurements of soot precursor (PAH) concentration, soot volume fractions [281, 372, 447, 527, 528], and through numerical simulations using detailed kinetic mechanisms [87, 380]. From these investigations, major differences in the sooting structures between SF and SFO flames can be identified, which are summarized as follows.

In a SFO flame, considerable amounts of OH radicals exist on the fuel side of the flame sheet, overlapping with the sooting zone. The spatial proximity of soot and OH species means that significant soot oxidation can take place as soot particles (formed on the fuel side of flame) are transported by the convective flow towards the high temperature flame sheet. In a SF flame, on the other hand, there is a large spatial separation between the flame sheet and the stagnation plane, and the convection is in a direction away from the flame. Consequently, unlike in SFO flames, oxidizing species like OH and O$_2$ do not coincide with the soot zone, so soot oxidation is absent [73]. These differences in soot evolution result in the distinct shape of SVF profiles between SFO and SF flames, as can be more clearly seen in Figure 11 [378]. In SFO flames, the profile of SVF is somewhat symmetric, indicating that the soot particles continuously experience inception, growth, and finally, destructive oxidation. The SVF drops to a negligible level before reaching the stagnation plane, consistent with the fact that SFO flames typically do not emit soot. In contrast, the SVF profile in SF flame is highly skewed toward the fuel side, characterized by a sharp decrease across the stagnation plane, a phenomenon observed also in several other studies [78, 281, 529, 530]. This is because the particles have very low diffusivity and cannot effectively penetrate through the stagnation plane, so they eventually leak out of the flame in the radial direction.

Of special note is the observation by D’Anna and coworkers [372, 531] that in a C$_2$H$_4$ SF flame ($X_{F,0} = 0.75$, $X_{O,0} = 0.22$ diluted by Ar), abundant small particles also exist on the fuel side of the
stagnation plane. The presence of soot in this pyrolytic region can be explained, according to the authors, by the formation of soot nuclei through physical coagulation of PAHs under low temperature conditions. At first glance, this observation seems to be incompatible from the measurements of Xu et al. [378], in which negligible amount of soot exists on the fuel side of particle stagnation plane (Figure 11). However, it is important to realize that D’Anna’s observation is consistent with the high soot number density on the fuel side of SF flames, as predicted by Xu et al. [378] through a detailed PAH-based soot model. The particles formed on the fuel side are not likely to experience significant mass-adding surface reactions due to the relatively low temperature in the region. Therefore, these particles are nascent and may have notably different optical properties compared to mature graphitic soot. They can even be liquid-like [18, 485] and transparent to visible/near infrared lights and so cannot be effectively detected by the visible or infrared light extinction method, as employed in the work of Xu et al. [378] and many others. On the other hand, these nanoparticles may well respond to the laser-induced emission spectrum method with the use of a UV laser (266 nm) as the excitation source, a technique frequently used by D’Anna and coworkers [372, 531].
Figure 11: Experimental and computed soot volume fraction profiles of C₂H₄ for (a) SFO flame (X_{F,0} = 0.28, X_{O,0} = 0.9) and (b) SF flame (X_{F,0} = 1.0, X_{O,0} = 0.3), reprinted from Xu et al. [378] with permission of Elsevier.

Another interesting difference between SF and SFO flames regards the profiles of PAH LIF signals, which were observed to have double peaks in SFO flames, but only one peak in SF flames [281, 447, 528]. Figure 12 shows a comparison of the measured SVF and PAH fluorescence profiles between typical C₂H₄ SFO (a) and SF (b) counterflow flames. In the SFO flames (a), a second increase of PAH fluorescence, starting at around z = 5mm, can be clearly seen. Considering that it occurs in the high-temperature soot oxidation zone, where soot oxidation by OH radicals leads to a decrease in SVF, Hwang et al. [281] tentatively attributed this second peak of LIF signal to PAHs generated by soot oxidative fragmentation.

Figure 12: Profiles of soot volume fraction, PAH fluorescence in SFO (a) and SF (b) ethylene CDFs. Also shown are computed C₂H₂, O₂ and OH mole fractions, reprinted from Hwang et al. [528] with permission of Elsevier.
Kang et al. [73] further investigated the sooting structure of normal \((X_{F,0}=1.0, X_{O,0}=0.21)\) and inverse coflow diffusion flames \((X_{F,0}=0.3, X_{O,0}=0.6)\) and observed a similarity in sooting characteristics between normal (inverse) coflow diffusion flame and SFO (SF) flame. This similarity is clearly demonstrated in Figure 13, where direct photos and sooting structures of these flames are presented. For the normal coflow flame (Figure 13a, central fuel jet surrounded by coflowing oxidizer), PAHs are formed in the fuel-rich pyrolysis zone near the fuel nozzle, and are then incepted to form soot particles, which are spatially distributed downstream of the PAH region. Soot particles are then transported further downstream towards the high temperature reaction zone, where highly oxidative OH radicals are abundant. Consequently, the axial profile of SVF and particle size both exhibited a rise-then-fall behavior with the increase in axial distance above the nozzle [62, 72, 323, 403, 532-534], similar to the case for a CDF of the SFO type.

For an inverse coflow flame (Figure 13b, central jet of oxidizer surrounded by coflowing fuel), the LIF and LII images obtained by Mikofski et al. [535] showed that the formation of PAHs and the soot zone occurred close to the flame wing, and the sooting zone almost completely enclosed the central reaction sheet—a fact that can also be visually confirmed by the relative position of the yellow luminous region and the blue flame zone in Figure 13d. As a result, after formation, soot particles are convected away from the high temperature flame sheet and are eventually emitted out from the flame. This evolution process is similar to that in SF CDFs; it also explains the experimental fact that the radially integrated SVF of inverse coflow increases and approaches a constant value downstream of the flame (i.e., no oxidation) [535, 536]. These studies demonstrated the important role of flow convection in the evolution of soot particles and thus in affecting the overall sooting structures of diffusion flames.
Figure 13. Sooting structures as visualized by OH LIF, PAH LIF and soot LII for normal (a) and inverse (b) coflow diffusion flames, reprinted from Mikofski et al. [535] with permission of Elsevier. Also shown are direct photos of normal coflow flame (c), reprinted from Bisetti et al. [537] with permission of Elsevier and inverse coflow flame (d), reprinted from Wang et al. [538] with permission of Elsevier.

4.2 Influences of stoichiometric mixture fraction on sooting structures in CDFs

The above discussions on the differences in soot evolution between SF and SFO flames highlight the important role of flame location in determining the sooting behavior in CDFs. The variation of flame location can be achieved by adjusting the dilution level in the fuel and/or oxidizer streams. The extent can be quantified by the stoichiometric mixture fraction, $Z_{st}$, defined as $Z_{st} = (1 + Y_{F,O}W_{O}/Y_{O,F}W_{F})^{-1}$. Here, $Y$ represents the mass fraction, $W$ is the molecular weight, $v_F$ and $v_O$ are fuel and oxygen stoichiometric coefficients, respectively. Therefore, the value of $Z_{st}$ will change when fuel or oxidizer compositions vary, indicating the change of flame structure. A typical SF flame with
undiluted fuel/air combustion (SFO flame with diluted fuel burning in pure oxygen) corresponds to a low (high) \( Z_{st} \) value.

Atreya et al. [539] investigated the effect of \( Z_{st} \) on soot formation in a series of low strain rate methane CDFs (effects of strain rate to be discussed in Section 4.3). The sooting structures of four flames, with \( Z_{st} \) increasing from 0.129 to 0.584, are compared in Figure 14. The data showed that as \( Z_{st} \) increased, the flame front (\( z_{T,\text{max}} \)) moved progressively towards the fuel side, along with the shift of the stagnation plane (\( z_{stg,p} \)) from the oxidizer to the fuel side. Such change of flame location notably affected the structures of the soot zone in many aspects. First, the flame with \( Z_{st} = 0.129 \) (a) had the largest soot zone thickness but the smallest average temperature in the soot zone; Second, SVF increased monotonically towards the stagnation plane (following the direction of particle convection) in flames with \( Z_{st} = 0.129 \) and 0.297 (a, b)—cases in which the reaction sheets were located on the oxidizer side of \( z_{stg,p} \). However, the SVF profiles exhibited a rise-then-fall behavior for flames with \( Z_{st} = 0.48 \) (c) and 0.584 (d), where the reaction front was either close to, or on the fuel side of, \( z_{stg,p} \). These results were consistent with the aforementioned sooting characteristics of typical SF and SFO flames; Third, particle number density (\( N \)) decreased continuously toward the stagnation plane for \( Z_{st} = 0.129 \), although it varied non-monotonically with \( z \) for the remaining three flames with larger \( Z_{st} \). This result suggests that the soot inception process can be strongly affected by the spatial shift of OH radicals towards the sooting zone with increasing \( Z_{st} \). As Atreya et al. [539] explained, this behavior was also compatible with the fact that the calculated OH mole fraction (without soot model) agreed well with the measured one for \( Z_{st} = 0.129 \), but obviously higher than the measured for cases with \( Z_{st} = 0.297 \) and 0.48—suggesting a fraction of OH radicals may be consumed for the oxidation of soot particles in the latter cases.
Figure 14. Flame and sooting zone structures for methane CDFs with varying $Z_{st}$. Data taken from Atreya et al. [539].

Also note in Figure 14, peak SVFs showed a non-monotonic variation with $Z_{st}$, with the flame of $Z_{st}=0.48$ having the highest peak SVF. This phenomenon is different from the observations in Refs. [377, 540-545], where peak SVF decreased monotonically with the increase in $Z_{st}$. Such differences may derive from the fact that the adiabatic flame temperatures remained unchanged as $Z_{st}$ varied in Refs. [377, 540-545], but they were not controlled by Atreya et al. [539]. With fixed peak temperature, results from Refs. [377, 540-545] generally showed that the inhibition of soot and their precursors with increasing $Z_{st}$ can be attributed to both the hydrodynamic effects and the alteration of flame structure.

The hydrodynamic effects refer to the directional change of soot convection when $Z_{st}$ changes [542]. The change in flame structure with $Z_{st}$ (i.e., the relationship between the temperature and local species concentration) may also affect soot formation significantly, as demonstrated by Du and Axelbaum.
through detailed experimental and numerical studies of ethylene CDFs. In their study, the sooting tendency of the CDFs decreased monotonically as $Z_{st}$ increased, attributed by the authors to the result of flame structure variation. Note, since only critical sooting condition was considered, the flames studied were only relevant to soot inception (i.e., without particle growth/transport processes), and therefore were expected not to be sensitive to the hydrodynamic effects. The general flame structures for flames with $Z_{st}$ of 0.064 and 0.78 are compared in Figure 15. As seen, the flame with higher $Z_{st}$ had lower fuel (marked as $Y_{C_2H_4}$) but higher oxygen concentrations ($Y_{O_2}$) near the reaction zone. Du and Axelbaum argued that in flames with high $Z_{st}$, the low fuel concentration tended to slow down fuel pyrolysis and soot precursor formation. In addition, the shift of oxygen concentration towards the fuel side enhanced the oxidation of soot precursors, also contributing to a reduction of soot formation. Similar arguments have been made by other researchers [377, 546, 547]. Furthermore, the sooting zone was known to be constrained in a region bounded by the oxidizing flame sheet and the fuel-rich zone, with temperatures high enough for soot inception [539, 540]. As $Z_{st}$ increased, the spatial shift of the oxidizing species towards the fuel nozzle reduced the thickness of the sooting zone until it reached a soot-free (permanently blue) condition [540].
Figure 15. Schematic diagram of flame structures for (a) ethylene/air CDFs with $Z_{st}$ of 0.064 and (b) diluted ethylene/oxygen CDF with $Z_{st}$ of 0.78, redrawn from Du et al. [540] with permission of Elsevier.

Through numerical simulations with detailed chemistry accounting for aromatic species formation, Skeen et al. [543] attributed the soot-free phenomena in high $Z_{st}$ flame to the inhibited formation of soot precursors (i.e., benzene and phenyl). In their series of ethylene CDFs investigated, an increase in $Z_{st}$ was seen to decrease the temperature in the fuel pyrolysis zone where abundant H radical existed, although the peak temperature remained constant. This, in turn, drove the dominant propargyl formation reactions (R50)–(R51) in the reverse direction because both (forward) reactions were endothermic and produced H radical. Consequently, the formation of benzene and phenyl via propargyl self-combination was inhibited.

\[
\begin{align*}
C_2H_2 + CH_3 & \leftrightarrow pC_3H_4 \text{ (propyne)} + H \quad \text{R50} \\
pC_3H_4 & \leftrightarrow C_3H_3 + H \quad \text{R51}
\end{align*}
\]

The change of $Z_{st}$ can result in flame structure variations in both physical distance and mixture fraction ($Z$) space, which may make it somewhat difficult to interpret the effects of $Z_{st}$ on sooting behavior. In such a context, Axelbaum and coworkers [77, 548] proposed to understand the flame structure effect in C/O ratio space. The parameter of C/O ratio can also indicate the local mixing process, and unlike the mixture fraction $Z$, it is insensitive to variation in fuel/oxidizer boundary composition. As shown in Figure 16a, when plotting the flame structure in terms of C/O ratio, the flame sheet is always located at (C/O)$_{st}$, which has a numerical value of 1/3 for ethylene fuel, regardless of $Z_{st}$. In contrast, the flame location changed notably with $Z_{st}$ when expressed in $Z$ space (see Figure 15). As an example of application, Xia et al. [77] analyzed the soot-free phenomena in high $Z_{st}$ flame reported by Skeen et al. [543]. Unlike the case where a considerable shift of H radical and temperature profiles occurred in the physical distance space [543], the edge of the H profile (fall-off) remained at nearly the same C/O ratio of 0.5 (Figure 16b). Therefore, it was straightforward to observe the reduction of flame temperature at C/O = 0.5 when $Z_{st}$ increased. In this regard, the
conclusion that it is the decrease of flame temperature in the H radical abundant region that causes the reduction of soot precursors in the high $Z_{st}$ flame, becomes more evident.

Figure 16: Global flame structure (a) and temperature and H radical profiles (b) of ethylene flames in C/O ratio space with different $Z_{st}$. Adapted from Xia et al [77] with permission of Elsevier.

4.3 Influences of strain rate on sooting structures in CDFs

Due to an accompanying significant entropy decrease [18], soot formations is a highly reversible and rate-limiting process; therefore it can be significantly affected by flow residence time. The characteristic flow time for flames under aerodynamic straining (i.e., CDF) is strain rate, which is essentially the gradient of axial velocity and has a unit of inverse time [549]. Considering that strain rate is a defining parameter for CDF, it is important to understand how it can affect sooting structures in CDFs.

Before discussing its effects, it is worthwhile to first clarify the definition of strain rate, since different researchers quantify this parameter in different ways. As a matter of fact, there is no true unique definition of strain rate in the counterflow, partly because of the axial variation of velocity.
gradient [550]. For studies with direct measurement of the axial flow field, either with LDV or PIV, it is a common practice to define the characteristic strain rate as the absolute value of the measured axial velocity gradient in the thermal mixing layer of the oxidizer stream [78, 462] (the oxidizer stream is chosen because most CDFs feature a very low stoichiometric mixture fraction so that the flame front is typically located on the oxidizer side of the stagnation plane). Figure 17 shows a typical axial velocity profile as measured by Du et al. [551]. The strain rate was found by fitting the experimental data with a straight line, as shown by the dashed line in the figure. This experimentally-determined strain rate will hereafter be referred to as the local strain rate ($K_L$). In the absence of experimental data, the strain rate may be estimated from nozzle exit velocities ($V_i$) and nozzle separation distances ($L$), where $i$ can be either fuel (F) or oxidizer (O) stream. The simplest approach is to assume a potential boundary condition and write the strain rate as $(V_F + V_O)/L$. This definition will hereafter be referred to as the global strain rate ($K_G$). It should be pointed out that the velocity boundaries of CDFs that employ nozzles with porous plugs are more like the plug flow type, and a more relevant description of the strain rate ($K_E$, effective strain rate) was provided by Seshadri and Williams [550, 552] as:

$$K_E = \frac{2V_O}{L} \left(1 + \frac{V_F}{V_O} \left(\frac{\rho_F}{\rho_O}\right)\right)$$  \hspace{1cm} \text{Eq. 23}

In the derivation of Eq. 23, in addition to employing a plug flow velocity boundary, it was further assumed that the viscous mixing layer could be treated as a thin sheet at the stagnation plane (i.e., large Reynolds number) [553]. Although experimentally determined $K_L$ is preferred, $K_G$ and $K_E$ also provide a convenient way to select corresponding experimental conditions for parametric studies on strain rate effects, as long as the definition is consistent. To avoid confusion when results from different studies are compared with different strain rate definitions, an explicit definition of which strain rate was employed will be offered, using the corresponding symbols. In the present section, only the effects of strain rate on overall sooting structure will be discussed; detailed parametric studies
will be reviewed in Section 5.5, where the unsteady response of soot formation to time-varying strain rates are also described.

**Figure 17.** Measured axial velocity profiles in counterflow diffusion flames and determination of local strain rate. Reprinted from Du et al. [551] with permission of Elsevier.

The strong effect of strain rate on sooting structures can be visualized in **Figure 18**, where the axial profiles of SVF, average particle size, and soot growth rate for a series of C$_2$H$_4$ CDFs ($X_{F,0}$=1.0, $X_{O,0}$=0.25) are shown [91]. With the nozzle exit velocity $V_0$ increasing from 15 cm/s to 30 cm/s, $K_E$ varied from 75 to 150 s$^{-1}$.

As can be seen, in addition to the decrease in peak SVF (a), the soot zone thickness decreased monotonically, consistent with the shrinkage in flame thickness as strain rate increased. Indeed, accompanying numerical simulation showed that with the increase in strain rate, the flame sheet and the stagnation plane were brought closer together, leaving an increasingly narrower region for soot growth. Average soot diameters also decreased (b), partially explained because the increased strain rate reduced time available for coagulation and surface growth between the soot inception zone and the particle stagnation plane. The volumetric soot mass growth rate $\omega V$ also decreased with the increase in strain rate, caused primarily by the reduced surface area in the more strained flames. However, the soot mass growth rate per unit surface area $\omega A$ showed an interesting behavior. In
regions close to the flame front $\omega_A$ was higher for flames with higher strain rate, while the trend became reversed in regions adjacent to the stagnation plane. The reason lies in the effects of strain rate on temperature profiles. Although the peak temperature was reduced due to flame weakening as the strain rate increased, in regions closer to the stagnation plane, where soot surface growth peaks, the flame with higher strain rate actually had a higher temperature. Nevertheless, the effects of this higher $\omega_A$ were overwhelmed by the lower surface area, a consequence of decreased PAHs, and thus, less soot nucleus, finally resulting in lower SVF in flames with higher strain rates.

![Diagram](image)

**Figure 18.** Effects of strain rate on axial profiles of soot volume fraction (a) average particle diameters, (b) soot mass growth rate, (c) and soot mass growth rate per unit area (d) in ethylene counterflow diffusion flames. Reprinted from Wang and Chung [91] with permission of Elsevier.

It is of interest to note here that the reduction of mass growth rate per unit surface area near the flame front, with an increase in straining, was also confirmed experimentally by Vandsburger et al.
In particular, the complementary measurement of the velocity profiles using LDV allowed the authors to determine the time derivative of soot concentrations, which—after being normalized by the total soot surface area as estimated from measured soot sizes—gives the soot surface growth rate.

Wang and Chung [91] concluded that the reduction of soot formation with increasing strain rate can be generally explained as: 1) reduced PAH concentration and soot inception rate, leading to less soot surface area per unit volume for mass growth to occur; (2) reduced soot surface growth rate per unit volume; and (3) reduced residence time (from inception region to particle stagnation plane) available for soot surface reactions to take place.

Before closing this subsection, we point out that for an originally-sooting CDF, progressively higher strain rates (shorter residence times) sequentially lead to the disappearance of soot, PAH and finally, extinction of the flame [551, 555]. In fact, the critical strain rate at which soot starts to disappear ($K_{cr}$) is frequently used as a quantitative indicator of the sooting tendency of CDFs.

4.4 Sooting structures of partially-premixed CDFs

In a counterflow configuration, various flame interaction modes can be realized, including single, double and triple flames [556-559], depending on the degrees of partial premixing in the fuel and/or oxidizer streams. Partially-premixed flames are especially interesting when it comes to soot formation, since the existence of oxygen in the fuel stream can notably influence soot formation through its effects on both flame structures and soot chemistry.

Mungekar and Atreya [560-562] investigated the effects of partial premixing of oxygen in the fuel stream on the sooting structures in methane CDFs for both SF and SFO configurations. Several interesting features were reported for SF flames [562]. First, all the partially-premixed flames investigated featured double flames, with the sooting zones sandwiched between the premixed (PF) and non-premixed (NP) reaction zones, as schematically shown in Figure 19 (b-d). Second, with an increase in the level of premixing (as the equivalence ratio in the fuel stream $\phi$ decreased from infinity
to $\phi = 3.4$), the diffusion flame front moved closer to the gas stagnation plane (marked as SP), because of fuel dilution and consumption by oxygen. In addition, the separation distance between the premixed and non-premixed flame front increased. This is because the rich premixed flame sheet tended to move to the fuel nozzle with the addition of oxygen, which can be explained by considering the balance between the stretched laminar flame speed and the incoming reactant velocity [563]. In particular, the flame speed of the premixture was increased due to oxygen addition so that the premixed flame sheet adjusted itself to a location with higher incoming reactant speed (i.e., closer to the fuel boundary). Third, the peak soot volume fraction ($f_v$) and the soot zone thickness decreased and then increased again with the increase in the level of partial premixing. As explained by the authors [561, 562], the initial decrease in soot zone thickness (from $\phi = \infty$ to 4.5) was due to the shift of the diffusion flame towards the gas stagnation plane; subsequent increase (from $\phi = 4.5$ to 3.4) was the result of increased spatial distance between the premixed and diffusion flame front, in which the soot zone was sandwiched. As to the variation of soot volume fraction, the authors attributed the initial reduction of $f_v$ to the dilution and chemical effects of oxygen addition. As more oxygen was added, however, the flame structure changed significantly because the location of peak temperature moved closer to the stagnation plane, resulting in increased temperature and longer soot residence time in the soot formation region. Consequently, peak $f_v$ increased again as $\phi$ decreased from 4.5 to 3.4. Compared to the case of $\phi = \infty$ (Figure 19a), a considerable amount of OH radicals existed in the soot zone for the partially-premixed case with $\phi = 3.4$. This led to the different sooting behavior: The soot particles leaked out near the stagnation plane for case of $\phi = \infty$, while they became oxidized when passing through the OH zone for $\phi = 3.4$.

For the SFO flames [561] with diffusion flame on the fuel side of the stagnation plane, partial premixing in the fuel side resulted in a monotonic reduction of SVF. These results again support the importance of flame structure changes on soot evolution.
In this section only the effect of partial-premixing on overall sooting structures is addressed. More detailed discussions on how partial premixing affects soot chemistry and quantitative soot loading will be discussed in a later section, when it will be seen that the effects can be drastically different for flames with different fuels.

Figure 19. Effects of oxygen addition on measured flame temperature, OH, and soot volume fraction $f_v$ for (a) purely diffusion CH$_4$ CDF (equivalence ratio in fuel stream $\phi = \infty$) and partially premixed cases with (b) $\phi = 6.0$, (c) 4.5 and (d) 3.4 (PF: premixed flame, SP: stagnation plane, NP: non-premixed flame). Data taken from Mungekar and Atreya [562].

5. Soot formation in counterflow diffusion flames: Parametric studies

Next follows a thorough review of parametric soot studies performed in counterflow diffusion flames. Through a comprehensive search of the literature, more than 100 articles have been identified and classified into six categories, based on the research objectives of these studies. These studies generally focused on examining various factors that affect soot formation and include the effects of 1) fuel types, 2) dilution, 3) pressure, 4) temperature, 5) strain rate, and 6) other parameters such as
radiation heat loss and electric field. For ease of reference Table 1 lists all relevant studies ordered by year of publication. For each individual study, fuel types, measured/predicted properties, diagnostic/numerical techniques, and the primary research objectives are listed. Note that several studies have more than one objective, such cases are grouped based on their primary objective of study, thus appearing only once in the list. Nevertheless, they can be referenced in different sub-sections in discussions that follow. Before examining the details, it is useful to summarize some general aspects of the factors that influence soot formation.

Generally, fuel molecular structures and additives determine pyrolysis products, thus affecting soot inception, surface growth and oxidation mainly through the influence on intermediate radical pools. As a controlling parameter for any kinetic process, temperature affects almost the entire soot formation process. Pressure and dilution influence soot inception and particle-particle coagulation by altering the concentration of important gaseous intermediates, as well as soot particles. Soot formation can also be significantly affected by flow field characteristics, primarily through their influence on flame structures, fuel/air mixing, strain rate and flame interactions. The electrical field may alter the flow field through ionic wind effects, also affecting soot formation, in addition to its potential role in altering the coagulation efficiency of charged soot particles.
Table 1: Summary of soot studies performed in counterflow flow flames

<table>
<thead>
<tr>
<th>Year/Author/Ref</th>
<th>Research Category</th>
<th>Fuel/Topic</th>
<th>Method</th>
</tr>
</thead>
</table>
| 1984, Vandsburger et al. [94] | Dilution effects | **Fuel:** Ethylene, propane  
**Topic:** Effects of oxygen concentration in the oxidizer stream on soot formation. | **EXP:** LDV, LE/LS |
| 1985, Vandsburger et al. [554] | Strain rate effect | **Fuel:** Ethylene  
**Topic:** Effects of oxidizer inlet velocity on soot formation of ethylene CDFs | **EXP:** LE/LS |
| 1987, Hura and Glassman [113] | Partial premixing | **Fuel:** Ethylene, propene, propane, iso-butane, n-butane  
**Topic:** Effect of oxygen as additive to various fuels on soot formation in CDF | **EXP:** LDV, LE/LS, LIF |
| 1988, Axelbaum et al. [74] | Fuel dilution; temperature effects | **Fuel:** Ethylene  
**Topic:** Separation of effects of flame temperature and fuel dilution on soot formation | **EXP:** LDV, LE/LS |
| 1989, Du et al. [551] | Fuel dilution; temperature effects | **Fuel:** Methane, ethane, propane, butane  
**Topic:** Separation of flame temperature and fuel dilution on soot formation for various fuels | **EXP:** LS, LIF |
| 1991, Du et al. [564] | Non-HC additive | **Fuel:** Ethylene, propane  
**Topic:** Effects of CO$_2$ and O$_2$ addition to fuel and oxidizer side on soot formation | **EXP:** LIF, LDV, LE |
| 1991, Leung, et al. [270] | Soot mechanism | **Fuel:** Ethylene, propane  
**Topic:** Phenomenological mechanism for soot formation, tested for ethylene and propane flames with varying oxygen concentrations in air streams | **SIM** |
| 1993, Baranov et al. [565] | Soot model | **Fuel:** Acetylene  
**Topic:** Soot model for spherical particles and soot aggregates in low pressure acetylene flames | **SIM** |
| 1993, Gomez and Rosner [79] | Sooting structure | **Fuel:** Methane  
**Topic:** Effect of thermophoresis on soot characteristics in methane counterflow diffusion methane flame | **EXP:** LE/LS, LDV |
| 1993, Taran [566] | Electric field | **Fuel:** Acetylene, benzene  
**Topic:** Soot formation in low pressure acetylene and benzene low flame subject to alternating electric fields | **EXP:** TEM |
| 1994, Delichatsios [567] | Soot model | **Fuel:** Ethylene, butadiene, butene, acetylene  
**Topic:** Phenomenological soot model for prediction of soot formation. | **SIM** |
| 1995, Du et al. [568] | Non-HC additive | **Fuel:** Ethylene, propane, butane  
**Topic:** Effects of CO, Ar, N$_2$, He, and H$_2$ as additives to ethylene, propane, and butane on soot formation | **EXP:** LS, LIF; |
<table>
<thead>
<tr>
<th>Year</th>
<th>Authors</th>
<th>Sooting/Process/Model</th>
<th>Fuel</th>
<th>Topic</th>
<th>EXP/SIM</th>
</tr>
</thead>
<tbody>
<tr>
<td>1995</td>
<td>Du et al. [76]</td>
<td>Sooting structure</td>
<td>Ethylene</td>
<td>Effects of stoichiometric mixture fraction on soot formation in counterflow and coflow flames</td>
<td>LDV SIM</td>
</tr>
<tr>
<td>1996</td>
<td>Tseng, et al. [570]</td>
<td>Partial premixing</td>
<td>Methane</td>
<td>Effects of oxygen addition in fuel on formation of gaseous intermediate</td>
<td>GC SIM</td>
</tr>
<tr>
<td>1996</td>
<td>Marquardt et al. [571]</td>
<td>Sooting process</td>
<td>Acetylene, carbon monoxide</td>
<td>Re-initiation of surface growth reactions in premixed counterflow flames</td>
<td>LE/LS SIM</td>
</tr>
<tr>
<td>1996</td>
<td>Wang et al. [78]</td>
<td>Soot model; strain rate effects</td>
<td>Ethylene</td>
<td>PAH-based soot model to study ethylene flames with different strain rates</td>
<td>LDV, LS/LE SIM</td>
</tr>
<tr>
<td>1997</td>
<td>Kang et al. [73]</td>
<td>Sooting structure</td>
<td>Propane</td>
<td>Effects of flow field on soot formation in counterflow, flames. Classification of SF and SFO flames.</td>
<td>LDV, LS/LE LIF</td>
</tr>
<tr>
<td>1997</td>
<td>Vincitore and Senkan [572]</td>
<td>Flame chemical structure</td>
<td>Methane</td>
<td>Major intermediate species and soot volume fraction profiles in methane CDF</td>
<td>GC/MS, LE/LS</td>
</tr>
<tr>
<td>1998</td>
<td>Chao et al. [544]</td>
<td>Sooting structure</td>
<td>Ethylene</td>
<td>Effects of hydrodynamics and transport characteristics on formation and oxidation of soot/soot precursors formation process</td>
<td>SIM</td>
</tr>
<tr>
<td>1998</td>
<td>Du et al. [573]</td>
<td>Pressure effect</td>
<td>Ethylene</td>
<td>Soot formation process in ethylene flames at various pressure (from 1 to 2.5 atm)</td>
<td>LS, LDV SIM</td>
</tr>
<tr>
<td>1998</td>
<td>Hwang et al. [165]</td>
<td>Partial premixing</td>
<td>Ethylene, propane</td>
<td>Effects of propane and oxygen addition soot formation in ethylene CDF</td>
<td>LDV, LE, LIF</td>
</tr>
<tr>
<td>1998</td>
<td>Hwang et al. [528]</td>
<td>Fuel mixing</td>
<td>Ethylene (propane)</td>
<td>Synergistic effect of soot formation processes with ethylene-propane mixtures.</td>
<td>LDV, LS/LE, LIF; SIM</td>
</tr>
<tr>
<td>1998</td>
<td>Sung et al. [85]</td>
<td>Pressure effect</td>
<td>Methane, propane</td>
<td>Sooting limits in methane and propane flames with pressures from 1 to 5 atm</td>
<td>Raman SIM</td>
</tr>
<tr>
<td>1998</td>
<td>Vincitore and Senkan [242]</td>
<td>Flame chemical structure; Fuel dilution</td>
<td>Ethane</td>
<td>Chemical (PAH in particular) speciation in ethane CDF with fuel dilution</td>
<td>GC/MS</td>
</tr>
<tr>
<td>1998</td>
<td>Olten and Senkan [407]</td>
<td>Flame chemical structure</td>
<td>Ethylene</td>
<td>Chemical speciation intermediates in ethylene counterflow flame</td>
<td>GC/MS</td>
</tr>
<tr>
<td>Year</td>
<td>Authors</td>
<td>Fuel mixing</td>
<td>Fuel: Propane (DMC, ethanol)</td>
<td>Topic: Effects of oxygenated additives (DMC and ethanol) on soot precursor formation</td>
<td>EXP: GC</td>
</tr>
<tr>
<td>------------</td>
<td>--------------------------</td>
<td>-------------</td>
<td>-----------------------------</td>
<td>-----------------------------------------------------------------------------------</td>
<td>---------</td>
</tr>
<tr>
<td>2000</td>
<td>Böhm and Lacas [575]</td>
<td>Pressure effect</td>
<td>Fuel: Methane</td>
<td>Topic: Effect of pressure (1 to 6 bar) on extinction limits and PAH formation in methane flames</td>
<td>EXP: UV LIF, SIM</td>
</tr>
<tr>
<td>2001</td>
<td>Beltrame et al. [576]</td>
<td>Dilution and strain rate effect</td>
<td>Fuel: Methane</td>
<td>Topic: Soot and NO formation in methane flames with various oxygen contents and strain rates (10-60s⁻¹)</td>
<td>EXP: LE SIM</td>
</tr>
<tr>
<td>2001</td>
<td>Böhm et al. [114]</td>
<td>Strain rate effect</td>
<td>Fuel: Methane</td>
<td>Topic: Soot precursor formation in methane CDF flames with various strain rates</td>
<td>EXP: LIF</td>
</tr>
<tr>
<td>2001</td>
<td>Santoianni et al. [577]</td>
<td>Strain rate effect</td>
<td>Fuel: Propane, methane</td>
<td>Topic: Soot formation in propane and methane flames subject to oscillating strain rates</td>
<td>EXP: LII PLIF, LDV, PIV</td>
</tr>
<tr>
<td>2002</td>
<td>Granata et al. [578]</td>
<td>Flame chemical structure</td>
<td>Fuel: 1,3-butadiene</td>
<td>Topic: Chemical speciation and kinetic modelling with emphasis on soot and PAH formation reaction pathways in 1,3-butadiene flame</td>
<td>EXP: GC/MS SIM</td>
</tr>
<tr>
<td>2003</td>
<td>Naik et al. [581]</td>
<td>Dilution effects</td>
<td>Fuel: Methane</td>
<td>Topic: Critical sooting conditions in terms of fuel/oxidizer dilution level</td>
<td>EXP: LIF SIM</td>
</tr>
<tr>
<td>2003</td>
<td>Violi et al. [87]</td>
<td>Soot model</td>
<td>Fuel: Ethylene</td>
<td>Topic: PAH-based kinetic modelling of soot formation in various ethylene diffusion flames</td>
<td>SIM</td>
</tr>
<tr>
<td>Year</td>
<td>Authors</td>
<td>Method</td>
<td>Fuel</td>
<td>Topic</td>
<td>Experiment</td>
</tr>
<tr>
<td>-----------</td>
<td>---------------------------------------------</td>
<td>-----------------</td>
<td>---------------</td>
<td>-----------------------------------------------------------------------</td>
<td>------------------</td>
</tr>
<tr>
<td>2004</td>
<td>Liu et al.</td>
<td>SIM</td>
<td>Ethylene</td>
<td>Effects of radiation heat loss on soot formation</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>2005</td>
<td>Berta et al.</td>
<td>EXP, GC, SIM</td>
<td>n-heptane</td>
<td>Effect of level of premixing on soot formation</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>2005</td>
<td>Kitajima et al.</td>
<td>EXP: GC/MS, LDV</td>
<td>Methane, propane</td>
<td>Effects of the dilution levels, flame temperature, and strain rates on PAH formation in methane and propane flames</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>2005</td>
<td>McNesby et al.</td>
<td>EXP: PLIF</td>
<td>Ethylene</td>
<td>Effect of ethanol addition on intermediate species and soot concentration</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>2005</td>
<td>Xiao et al.</td>
<td>EXP: PLIF</td>
<td>Methane, propane</td>
<td>Effect of strain rate on PAH and soot formation in steady and unsteady CDFs</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>2005</td>
<td>Yoon et al.</td>
<td>EXP: PLII, LIF, SIM</td>
<td>Ethylene</td>
<td>Kinetic effects of methane, ethane, propane, and propene mixing on PAH and soot formation</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>2006</td>
<td>Berta et al.</td>
<td>EXP: GC, SIM</td>
<td>n-heptane</td>
<td>NO\textsubscript{x} and PAH formation in n-heptane counter-flow diffusion flame with various partial premixing levels</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>2006</td>
<td>Mungekar and Atreya</td>
<td>EXP: LE/LS, TEM, LIF SIM</td>
<td>Methane</td>
<td>Effects of premixing level on radiation and sooting characteristics</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>2006</td>
<td>Mungekar and Atreya</td>
<td>EXP: TEM, LIF, LE/LS SIM</td>
<td>Methane</td>
<td>Flame radiation and soot formation in methane flames with fuel-side oxygen addition</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>2007</td>
<td>Chung and Violi</td>
<td>SIM</td>
<td>Propene</td>
<td>Importance of oxidative reactions on soot growth and the dependence of soot structure on combustion conditions</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>2007</td>
<td>Li et al.</td>
<td>EXP: SEM</td>
<td>Methane</td>
<td>Effects of catalyst, temperature, and strain rate of the flow on carbon nanotubes synthesis.</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>2007</td>
<td>Sarathy et al.</td>
<td>EXP: GC</td>
<td>Methyl- crotonate, methyl-butanoate</td>
<td>Effects of the FAME molecular structure on the formation of intermediate species.</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Year</td>
<td>Authors</td>
<td>Fuel</td>
<td>Topic</td>
<td>Method(s)</td>
<td></td>
</tr>
<tr>
<td>--------</td>
<td>---------------------------------</td>
<td>-----------------------</td>
<td>----------------------------------------------------------------------</td>
<td>------------</td>
<td></td>
</tr>
<tr>
<td>2007</td>
<td>Yamamoto et al. [590]</td>
<td>Acetylene</td>
<td>Effects of strain rate on PAH formation in CDFs</td>
<td>GC/MS</td>
<td></td>
</tr>
<tr>
<td>2007</td>
<td>Yoo and Im [591]</td>
<td>Ethylene</td>
<td>Soot dynamics in turbulent counterflow flames</td>
<td>SIM</td>
<td></td>
</tr>
<tr>
<td>2008</td>
<td>Berta et al. [592]</td>
<td>n-Heptane</td>
<td>Formation pathways for aromatic species</td>
<td>EXP: GC</td>
<td></td>
</tr>
<tr>
<td>2008</td>
<td>Cuoci et al. [593]</td>
<td>Methane</td>
<td>Frequency response of PAH concentration to strain rate harmonic oscillations</td>
<td>SIM</td>
<td></td>
</tr>
<tr>
<td>2008</td>
<td>Leylegian et al. [594]</td>
<td>Methane, Ethylene</td>
<td>Effects of chloromethane addition on soot formation</td>
<td>EXP: LDV, LS, SIM</td>
<td></td>
</tr>
<tr>
<td>2008</td>
<td>Yoon et al. [100]</td>
<td>Methane, ethane, propane, ethylene</td>
<td>Effect of the addition of dimethyl ether on polycyclic aromatic hydrocarbon and soot formation</td>
<td>EXP: LII, LIF, SIM</td>
<td></td>
</tr>
<tr>
<td>2008</td>
<td>Zhong and Xi [595]</td>
<td>n-heptane</td>
<td>Reduction of a detailed n-heptane chemical kinetic mechanism (validated in CDF)</td>
<td>SIM</td>
<td></td>
</tr>
<tr>
<td>2009</td>
<td>Blanquart et al. [596]</td>
<td>Acetylene, n-heptane</td>
<td>Development of a kinetic model with PAH formation (validated in CDF)</td>
<td>SIM</td>
<td></td>
</tr>
<tr>
<td>2009</td>
<td>Cuoci et al. [395]</td>
<td>Propane</td>
<td>Response of soot volume fraction to strain rate oscillations</td>
<td>SIM</td>
<td></td>
</tr>
<tr>
<td>2009</td>
<td>Cuoci et al. [598]</td>
<td>Methane, propane, ethylene</td>
<td>PAH and soot formation in unsteady counterflow diffusion flames with strain rate oscillations</td>
<td>SIM</td>
<td></td>
</tr>
<tr>
<td>2009</td>
<td>D'Anna et al. [531]</td>
<td>Ethylene</td>
<td>Soot growth and evolution in ethylene CDF</td>
<td>EXP: LII, LIF, SIM</td>
<td></td>
</tr>
<tr>
<td>2009</td>
<td>Hou et al. [599]</td>
<td>Ethylene, methane</td>
<td>Effects of fuel/oxygen concentration on carbon nano-onions formation</td>
<td>EXP: SEM, TEM</td>
<td></td>
</tr>
<tr>
<td>2009</td>
<td>Jahangirian [80]</td>
<td>Ethylene</td>
<td>Soot precursor formation in ethylene CDF doped with jet surrogate fuels</td>
<td>EXP: GC/MS</td>
<td></td>
</tr>
<tr>
<td>2009</td>
<td>Mehta et al. [600]</td>
<td>Ethylene</td>
<td>Sensitivity of soot model parameters in modeling soot in CDF.</td>
<td>SIM</td>
<td></td>
</tr>
<tr>
<td>Year</td>
<td>Authors</td>
<td>Experiment Type</td>
<td>Fuel Type</td>
<td>Topic</td>
<td>Reference</td>
</tr>
<tr>
<td>--------</td>
<td>--------------------------</td>
<td>-----------------</td>
<td>-----------------------------</td>
<td>-------------------------------------------------------------------------------------------------</td>
<td>-----------</td>
</tr>
<tr>
<td>2010</td>
<td>Frassoldati et al. [601]</td>
<td></td>
<td><strong>Fuel:</strong> n-propanol, iso-propanol</td>
<td><strong>Topic:</strong> Chemical speciation and kinetic analysis of n-propanol and iso-propanol flames, with discussions on soot precursors.</td>
<td>EXP: GC/MS SIM</td>
</tr>
<tr>
<td>2010, Karatas et al. [86]</td>
<td>Fuel mixing</td>
<td><strong>Fuel:</strong> n-butane, iso-butane, ethylene, propane</td>
<td><strong>Topic:</strong> Synergistic effect of fuel mixture on soot formation</td>
<td></td>
<td>EXP:</td>
</tr>
<tr>
<td>2011, Choi et al. [447]</td>
<td>Fuel mixing</td>
<td><strong>Fuel:</strong> n-heptane, iso-octane, toluene</td>
<td><strong>Topic:</strong> Synergistic effect on PAH formation for n-heptane/toluene and iso-octane/toluene mixtures</td>
<td></td>
<td>EXP: LIF, LII SIM</td>
</tr>
<tr>
<td>2011, Choi et al. [602]</td>
<td>Fuel mixing</td>
<td><strong>Fuel:</strong> Ethylene</td>
<td><strong>Topic:</strong> Kinetic insights on the effects of methane, ethane, or propane addition in ethylene fuels on soot formation</td>
<td></td>
<td>EXP: LE/LS SIM</td>
</tr>
<tr>
<td>2011, Hayashi et al. [603]</td>
<td>Spray flame</td>
<td><strong>Fuel:</strong> n-decane</td>
<td><strong>Topic:</strong> Effects of fuel droplet size on soot formation in spray flames</td>
<td></td>
<td>EXP: LII, TR-LII; SIM</td>
</tr>
<tr>
<td>2011, Narayanan et al. [604]</td>
<td>Soot radiation</td>
<td><strong>Fuel:</strong> Ethylene</td>
<td><strong>Topic:</strong> Effect of radiation heat loss from soot particles on flame extinction</td>
<td></td>
<td>SIM</td>
</tr>
<tr>
<td>2011, Seiser et al. [605]</td>
<td>Fuel type</td>
<td><strong>Fuel:</strong> n-decane, JP-10</td>
<td><strong>Topic:</strong> Chemical speciation and kinetic modelling of n-decane and JP-10 flames, with discussions on soot precursors</td>
<td></td>
<td>EXP: GC SIM</td>
</tr>
<tr>
<td>2012, Carbone and Gomez [606]</td>
<td>Fuel mixing</td>
<td><strong>Fuel:</strong> Methane, ethylene (toluene)</td>
<td><strong>Topic:</strong> Chemical structures of toluene-doped methane flame and ethylene flame</td>
<td></td>
<td>EXP: GC/MS SIM</td>
</tr>
<tr>
<td>2012, Feng et al. [82]</td>
<td>Fuel type</td>
<td><strong>Fuel:</strong> Biodiesel</td>
<td><strong>Topic:</strong> Sooting tendencies of various biodiesel fuels</td>
<td></td>
<td>EXP: GC/MS, LE SIM</td>
</tr>
<tr>
<td>2012, Raj et al. [203]</td>
<td>Kinetic mechanism</td>
<td><strong>Fuel:</strong> iso-octane, n-heptane, toluene</td>
<td><strong>Topic:</strong> Mechanism development for gasoline surrogate fuels with an emphasis on the formation of large PAHs</td>
<td></td>
<td>SIM</td>
</tr>
<tr>
<td>2012, Salamanca et al. [607]</td>
<td>Fuel mixing</td>
<td><strong>Fuel:</strong> Ethylene</td>
<td><strong>Topic:</strong> Effect of ethanol addition on particulate formation in ethylene flames</td>
<td></td>
<td>EXP: LIF, LII, TR-LIE</td>
</tr>
<tr>
<td>2012, Sirignano et al. [608]</td>
<td>Sooting structure</td>
<td><strong>Fuel:</strong> Ethylene</td>
<td><strong>Topic:</strong> Demonstrating the correlation of fluorescence emission spectra and fluorescence lifetimes with the characteristics of the PAH/incipient soot particles</td>
<td></td>
<td>EXP: TR-LII, LIF</td>
</tr>
<tr>
<td>2012, Slavinskaya et al. [63]</td>
<td>Kinetic mechanism</td>
<td><strong>Fuel:</strong> Ethylene, ethane</td>
<td><strong>Topic:</strong> Development of PAH growth chemical kinetic mechanism with validation in CDF</td>
<td></td>
<td>SIM</td>
</tr>
<tr>
<td>2013, Carbone and Gomez [609]</td>
<td>Fuel mixing</td>
<td><strong>Fuel:</strong> Methane, ethylene (1,2,4-trimethyl benzene)</td>
<td><strong>Topic:</strong> Chemical effect of 1,2,4-trimethyl benzene addition in Methane/ethylene flames</td>
<td></td>
<td>EXP: GC/MS SIM</td>
</tr>
<tr>
<td>Year</td>
<td>Authors</td>
<td>Type</td>
<td>Fuel</td>
<td>Topic</td>
<td>Methodology</td>
</tr>
<tr>
<td>--------------</td>
<td>----------------------</td>
<td>-----------------------</td>
<td>------</td>
<td>----------------------------------------------------------------------</td>
<td>-------------</td>
</tr>
<tr>
<td>2013, Han, et al. [611]</td>
<td>Fuel type</td>
<td>Fuel: n-heptane, 1-heptene</td>
<td>Topic: Effect of the unsaturated bond in the fuel molecular structure on NOx and soot emissions lean premixed, rich premixed, and non-premixed combustion</td>
<td></td>
<td></td>
</tr>
<tr>
<td>2013, Hayashi et al. [612]</td>
<td>Spray flame</td>
<td>Fuel: n-decane</td>
<td>Topic: Effects of fuel droplet size distribution on soot formation</td>
<td></td>
<td></td>
</tr>
<tr>
<td>2013, Hernández et al. [613]</td>
<td>Soot radiation</td>
<td>Fuel: Ethylene</td>
<td>Topic: Effects of soot and gas radiation on soot formation and flame structure</td>
<td></td>
<td></td>
</tr>
<tr>
<td>2013, Skeen et al. [615]</td>
<td>Soot chemical properties</td>
<td>Fuel: Acetylene, ethylene, propane</td>
<td>Topic: Chemical identification of species condensed on soot particles</td>
<td></td>
<td></td>
</tr>
<tr>
<td>2013, Skeen et al. [413]</td>
<td>Pressure effects</td>
<td>Fuel: Acetylene</td>
<td>Topic: Chemical speciation composition of acetylene flames at various pressures</td>
<td></td>
<td></td>
</tr>
<tr>
<td>2013, Xia et al. [77]</td>
<td>Sooting structure</td>
<td>Fuel: Ethylene, propane</td>
<td>Topic: utility of C/O ratio space in interpreting the effect of $Z_{st}$ on sooting structure</td>
<td></td>
<td></td>
</tr>
<tr>
<td>2014, Figura and Gomez [618]</td>
<td>Pressure effects</td>
<td>Fuel: Ethylene</td>
<td>Topic: Chemical structure of ethylene flames at pressures up to 2.5 Mpa (0.1-2.5Mpa) with discussions on soot precursors</td>
<td></td>
<td></td>
</tr>
<tr>
<td>2014, Park et al. [527]</td>
<td>Electrical fields</td>
<td>Fuel: Ethylene</td>
<td>Topic: Effect of DC electric field on soot reduction</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Year</td>
<td>Authors</td>
<td>Topic</td>
<td>Fuel Type</td>
<td>Fuel Additive</td>
<td>Other Details</td>
</tr>
<tr>
<td>-----------</td>
<td>-----------------------------</td>
<td>------------------------------------------------------------------------</td>
<td>---------------------------</td>
<td>---------------</td>
<td>--------------------------------------------------------------------------------</td>
</tr>
<tr>
<td>2015</td>
<td>Carbone et al. [81]</td>
<td>Development of CDF-based quantitative indices of sooting tendencies</td>
<td>Ethylene</td>
<td></td>
<td>EXP: GC/MS, SIM:</td>
</tr>
<tr>
<td>2015</td>
<td>Carbone and Gomez [621]</td>
<td>Fuel mixing</td>
<td>Methane, ethylene</td>
<td></td>
<td>EXP: GC/MS, SIM:</td>
</tr>
<tr>
<td>2015</td>
<td>Choi et al. [476]</td>
<td>Fuel mixing</td>
<td>Ethylene (DME)</td>
<td></td>
<td>EXP: LE/LS, SIM:</td>
</tr>
<tr>
<td>2015</td>
<td>Choi et al. [622]</td>
<td>Fuel mixing</td>
<td>Ethylene</td>
<td></td>
<td>EXP: LIF, LII, SIM:</td>
</tr>
<tr>
<td>2015</td>
<td>Figura et al. [623]</td>
<td>Fuel mixing</td>
<td>Ethylene</td>
<td></td>
<td>EXP: GC/MS</td>
</tr>
<tr>
<td>2015</td>
<td>Johansson et al. [136]</td>
<td>Kinetic pathway</td>
<td>Ethylene</td>
<td></td>
<td>EXP: TOF-MS, SIM:</td>
</tr>
<tr>
<td>2015</td>
<td>Raj et al. [261]</td>
<td>Fuel additive</td>
<td>Methane</td>
<td></td>
<td>EXP: GC, FTIR, XRD</td>
</tr>
<tr>
<td>2015</td>
<td>Schenk et al. [208]</td>
<td>Fuel type, kinetic pathway</td>
<td>n-butane, i-butane, i-butene, i-butanol</td>
<td></td>
<td>EXP: GC/MS, Helium-ion microscopy</td>
</tr>
<tr>
<td>2015</td>
<td>Sirignano et al. [372]</td>
<td>Sooting structure; kinetic pathway</td>
<td>Ethylene</td>
<td></td>
<td>EXP: LII, LS</td>
</tr>
<tr>
<td>2015</td>
<td>Wang et al. [140]</td>
<td>Soot model; fuel mixing</td>
<td>Ethylene</td>
<td></td>
<td>EXP: LE/LS, SIM</td>
</tr>
<tr>
<td>2016</td>
<td>Choi et al. [624]</td>
<td>Non-HC additive</td>
<td>Ethylene</td>
<td></td>
<td>EXP: TEM</td>
</tr>
<tr>
<td>2016</td>
<td>Selvaraj et al. [625]</td>
<td>Soot model</td>
<td>Ethylene</td>
<td></td>
<td>SIM</td>
</tr>
<tr>
<td>Year, Authors [Ref]</td>
<td>Fuel Type; Fuel Mixing</td>
<td>Topic</td>
<td>EXP; SIM</td>
<td></td>
<td></td>
</tr>
<tr>
<td>---------------------</td>
<td>------------------------</td>
<td>-------</td>
<td>---------</td>
<td></td>
<td></td>
</tr>
<tr>
<td>2016, Zimmer et al. [627]</td>
<td>Fuel: Ethylene</td>
<td>Effects of mass and energy coupling between the gas- and the solid-phase.</td>
<td>SIM</td>
<td></td>
<td></td>
</tr>
<tr>
<td>2017, Carbone et al. [628]</td>
<td>Fuel: Ethylene</td>
<td>Chemical speciation of incipiently sooting partially premixed flames of ethylene at various pressures</td>
<td>GC/MS SIM</td>
<td></td>
<td></td>
</tr>
<tr>
<td>2017, Hansen et al. [207]</td>
<td>Fuel: n-butane, i-butane, i-butene</td>
<td>Kinetic pathways for PAH formation in flames of C4 fuels</td>
<td>MBMS SIM</td>
<td></td>
<td></td>
</tr>
<tr>
<td>2017, Park et al. [629]</td>
<td>Fuel: n-heptane, iso-octane, toluene</td>
<td>Compositional effects on the sooting tendencies of toluene primary reference fuels</td>
<td>LII, LE SIM</td>
<td></td>
<td></td>
</tr>
<tr>
<td>2017, Rodrigues et al. [630]</td>
<td>Fuel: Propane</td>
<td>Sooting and PAH formation characteristics of CDF subject to strain rate harmonic oscillations</td>
<td>SIM</td>
<td></td>
<td></td>
</tr>
<tr>
<td>2017, Xue et al. [631]</td>
<td>Fuel: Jet fuels</td>
<td>Sooting tendencies of representative conventional and alternative jet fuels</td>
<td>LII, LE SIM</td>
<td></td>
<td></td>
</tr>
<tr>
<td>2018, Amin et al. [632]</td>
<td>Fuel: Ethylene</td>
<td>Effects of pressure on primary particle size distribution and fractal properties of soot aggregates</td>
<td>TEM SIM</td>
<td></td>
<td></td>
</tr>
<tr>
<td>2018, Gleason et al. [489]</td>
<td>Fuel: Ethylene</td>
<td>Effects of temperature on soot evolution and optical properties in highly controlled CDFs</td>
<td>pyrometry SIM</td>
<td></td>
<td></td>
</tr>
<tr>
<td>2018, Kalvakala et al. [377]</td>
<td>Fuel: Ethylene, propane, propene</td>
<td>Effects of fuel unsaturation and stoichiometric mixture fraction on NOx, PAH and soot formation</td>
<td>SIM</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Year</td>
<td>Author(s)</td>
<td>Type</td>
<td>Fuel</td>
<td>Topic</td>
<td>Experimentation</td>
</tr>
<tr>
<td>------------</td>
<td>--------------------</td>
<td>----------------------</td>
<td>-------------------------------</td>
<td>----------------------------------------------------------------------</td>
<td>-----------------</td>
</tr>
<tr>
<td>2018, Li</td>
<td>[633]</td>
<td>Fuel mixing; pressure effects</td>
<td>Ethylene</td>
<td>Effects of DME addition to ethylene on sooting limits at elevated pressures</td>
<td>EXP: LS SIM</td>
</tr>
<tr>
<td>2018, Raj et al.</td>
<td>[634]</td>
<td>Fuel additive</td>
<td>Methane</td>
<td>Effects of iron pentacarbonyl addition on soot formation and oxidation in methane CDF</td>
<td>EXP: SMPS SEM</td>
</tr>
<tr>
<td>2018, Sarnacki and Chelliah</td>
<td>[530]</td>
<td>Strain rate effects; pressure effects</td>
<td>Ethylene</td>
<td>Effects of strain rate and pressure (up to 30 atm) on soot formation</td>
<td>EXP: LII, PIV</td>
</tr>
<tr>
<td>2018, Xu et al.</td>
<td>[378]</td>
<td>Dilution effects; flame structure effects</td>
<td>Ethylene</td>
<td>Kinetic analysis on the effects of oxidizer composition on soot formation in both SF and SFO flames</td>
<td>EXP: LE SIM</td>
</tr>
<tr>
<td>2018, Xue et al.</td>
<td>[636]</td>
<td>Pressure effects</td>
<td>Ethylene</td>
<td>Pressure effect on soot formation and the implications of keeping density-weighted strain rate constant</td>
<td>EXP: LII SIM</td>
</tr>
<tr>
<td>2019, Yan et al.</td>
<td>[638]</td>
<td>Soot evolution process</td>
<td>Ethylene</td>
<td>Dependence of wavelength on soot extinction measurements</td>
<td>EXP: LE</td>
</tr>
<tr>
<td>2019, Gleason et al.</td>
<td>[490]</td>
<td>Temperature effects; pressure effects</td>
<td>Ethylene</td>
<td>Effects of temperature and pressure on soot evolution and optical properties in highly controlled CDFs</td>
<td>EXP: pyrometry</td>
</tr>
<tr>
<td>2019, Yan et al.</td>
<td>[397]</td>
<td>Fuel mixing</td>
<td>Ethylene</td>
<td>Effects of methanol and ethanol addition on soot formation in ethylene CDF</td>
<td>EXP: LII, LE SIM</td>
</tr>
<tr>
<td>2019, Hoerlle and Pereira</td>
<td>[639]</td>
<td>Dilution effect</td>
<td>Ethylene</td>
<td>Effects of CO2 addition in oxygen-enriched CDF of ethylene</td>
<td>SIM</td>
</tr>
<tr>
<td>2019, Mahmoud et al.</td>
<td>[640]</td>
<td>Dilution effect</td>
<td>Ethylene</td>
<td>Combined effects of CO2 and H2O addition in fuel- and oxidizer-side of ethylene CDF</td>
<td>EXP: LII, LE SIM</td>
</tr>
</tbody>
</table>
5.1 Effects of fuel types on soot formation

It is natural to imagine that certain fuels are inherently more sooting than others, and it is generally accepted that fuel molecular structure plays a vital role in determining the sooting tendency of a combustion system [96, 641]. This point could be best viewed from the fact that flames fueled with different isomers may exhibit significantly different sooting characteristics, although they share similar flame temperature, C/O ratio, and other flow field parameters. For instance, in counterflow flames of neat fuels and synthetic air ($X_{O,0} = 0.25$), it was observed that the soot volume fraction of iso-butane flame was three times as much as that of n-butane flame, although the two flames had nearly the same peak temperatures [113].

Fundamental understanding of the effect of fuel molecular structure on soot formation requires comprehensive knowledge of the kinetic pathways from the fuel to the molecular soot precursors, as well as the interactions between the pool of intermediate species and the soot surface growth. An important way to obtain such information is through numerical simulations with a detailed chemistry. Because of its simple quasi one-dimensional nature of flow field, the CDF configuration is particularly amenable to such detailed computation. In general, there are two types of studies that focus on fuel effects. Flames can either be established using neat fuels of the target [635] or various target fuels can be doped into a baseline flame of—for example—ethylene [642, 643] or methane [67].

Before discussing individual investigations, it is important to mention here that studies comparing the sooting tendency of different fuels should be considered in view of the changes in other parameters (e.g. stoichiometric mixture fraction and flame maximum/adiabatic temperature) that occur when using different fuels. In fact, it is always important to keep in mind that the conclusion that we draw from a specific parametric study must be understood with the effects of accompanying variations of other parameters taken into full consideration.
5.1.1 Counterflow diffusion flames with single component (neat) fuels

Glassman and coworkers are among the pioneers using CDFs to study the fuel-dependence on soot formation [94, 113]. These relatively early studies utilized the configuration in which a flame is established in the forward stagnation region of a cylindrical porous burner, as schematically shown in Figure 20(a). In Ref. [113], a uniform flow of synthetic air ($X_{O,0} = 0.25$) was set to approach the porous burner with a free-stream velocity of 30 cm/s. The measurements were made along the centerline for axial velocity, temperature, and soot volume fraction/size/number density using LDV, thermocouples, and light extinction/scattering techniques, respectively. Propane, n-butane, and iso-butane fuels were tested, and the soot-related results are summarized in Figure 20 (b). Note that in the original work [113], additional flames of alkene fuels such as ethylene and propene were also established; however, the oxidizer concentrations ($X_{O,0}$) used were different than those used for alkane fuels, making a direct comparison of sooting tendencies across these different fuel classes difficult. Therefore, only the results for alkane fuels are re-plotted in Figure 20.
Figure 20. Schematic of counterflow flow diffusion flame stabilized around a porous cylindrical burner (a), redrawn from Hura and Glassman [113]; Measured soot volume fraction (b), average particle size (c), and number density (d) as a function of time from the flame front towards the burner, data taken from Hura and Glassman [113].

As can be seen, the iso-butane flame generated the highest level of soot, followed by propane and n-butane (b). In addition, the higher sooting fuels also produced larger soot particles (c). However, the number density profiles (d) were confined in a relatively narrow band and exhibited less variation among the different fuels, believed to be consistent with the behavior of a coagulation-dominated aerosol system. Due to the lack of well-validated soot mechanisms that were able to handle multiple fuels in the early years, the authors did not attempt to explain these experimentally-observed trends through direct soot simulations of the CDF. Instead, inferences were taken from relevant iso-thermal
pyrolysis studies and it was suggested that the sooting tendency was dependent not only on fuel pyrolysis rates but also on the intermediate species formed during the pyrolysis process [96, 113, 644]. This observation, regarding iso-butane’s higher sooting tendency as compared to n-butane, was consistent with an earlier ranking using the smoke point data [645]. It was also corroborated by a number of subsequent studies [86, 231, 620]. In particular, it was pointed out that the decomposition of iso-butane leads to rather high levels of propene and iso-butene [646], which can be more readily converted to propargyl, and subsequently to benzene [152], as compared to C₂ species, which are among the major decomposition products of n-butane [647]. An exception comes from the work of Singh et al. [84], where it was observed that the peak soot volume fraction of the n-butane CDF was notably higher than that of iso-butane flame under the same conditions of \( X_{F,0} = 0.29, X_{O,0} = 0.9 \), and effective strain rate \( K_E = 57 \text{ s}^{-1} \). As an attempt to rationalize their experimental results, the authors performed kinetic simulations of the respective flames using both the Merchant [648] and the USC II mechanisms [649]. The numerical results from the two mechanisms consistently showed that although n-butane flame had a higher concentration of C\(_2\)H\(_2\), it featured a notably lower level of C\(_3\)H\(_3\) than iso-butane flames. This is consistent with our previous discussions on the decomposition products of the two butane isomers. Further considering the importance of the propargyl recombination reaction in forming aromatic species, it was expected (and later confirmed by PAH LIF experiments [441]) that iso-butane would produce more aromatic soot precursors. These findings allowed the authors to conclude that the ranking of soot tendencies may not be consistent with concentrations of aromatics in the inception stage [84, 441]. However, further research is still needed to clarify the details of how the trends became reversed during soot growth processes.

Using an opposed-nozzle counterflow configuration, Wang and Chung [620] performed experimental work on the sooting limit to rank the sooting tendencies of various C\(_1\)–C\(_4\) gaseous hydrocarbons. The results showed that for SF flames, propane and n-butane have comparable sooting tendencies, which are notably higher than those of methane and ethane. In addition, the soot formation
with C₂ and C₃ alkene fuels was significantly higher than ethane and propane, respectively, and the branched iso-butane was more sooting than its normal-chain counterparts. Kalvakala et al. [377] computationally investigated the effects of fuel unsaturation on soot formation by comparing the predicted peak soot volume fraction among counterflow diffusion flames of ethylene, propane, and propene. In agreement with the experimental results of Wang and Chung [620], they observed that the computed sooting tendency follows propene > ethylene > propane, regardless of the stoichiometric mixture fractions at which the comparisons were made [377]. Accompanying kinetic simulations conducted at \( X_{F,0} = 1.0 \) and \( X_{O,0} = 0.21 \) demonstrated that the major benzene formation pathway was similar in both propane and propene flames, being the recombination of propargyl radicals. As such, their different sooting tendency was largely a result of the different capability of the respective fuel pyrolysis products in forming propargyl radicals. Due to the presence of the double bond, the C-H bond of the γ carbon had the lowest bond energy, so that a major decomposition product of propene is allyl radical (\( \text{a-C}_3\text{H}_5 \)), which through multiple hydrogen abstractions can be readily converted to produce \( \text{C}_3\text{H}_3 \). However, for propane (\( \text{C}_3\text{H}_8 \)) molecules, the C-C bond was relatively weaker than the C-H bond, so that ethyl radical became the most dominant decomposition product, while \( \text{C}_3 \) species like n-propyl and propene became secondary. The fact that propane was less sooting than propene was then rationalized by noting that it is significantly more difficult to form \( \text{C}_3\text{H}_3 \) from the main decomposition products of \( \text{C}_3\text{H}_8 \) (i.e., \( \text{C}_2 \) species) than from unsaturated \( \text{C}_3 \) species.

The effects of unsaturation on PAH/soot formation were also investigated for \( \text{C}_7 \) hydrocarbon fuels (n-heptane and 1-heptene) in CDFs [610, 611]. Utilizing a detailed reaction mechanism and a PAH-based soot model, Fu et al. [610] showed that 1-heptene flame has a consistently higher soot number density, average particle size, and soot volume fraction as compared to the n-heptane flame, although both flames had very similar heat release characteristics. Furthermore, the numerical results also indicated that the existence of the double bond exerted increasingly pronounced effects on PAH and soot formation as the flame strain rates and/or levels of fuel stream partial premixing increased.
Alkene fuels are important constituents of practical hydrocarbon fuels [650, 651]; they are also key intermediates during alkane oxidation and pyrolysis [652]. As such, detailed knowledge of alkene combustion is required if the combustion and pollutant formation mechanisms of more practical fuels are to be understood. Motivated by this, Wang et al. [635] performed a systematic investigation on the sooting tendencies of C$_2$-C$_8$ 1-alkene fuels in CDFs. The experimental results from the LII and PAH LIF measurements, summarized in Figure 21, indicated that 1-butene (ethylene) flame produced most (least) soot and PAHs, and the sooting tendency decreased with the increase in the number of carbon atoms for fuels heavier than 1-C$_4$H$_8$. Accompanying numerical simulation confirmed and provided kinetic insights into these experimental trends.

![Figure 21](image.png)

Figure 21. Experimental peak PAH LIF signals detected at various wavelengths and peak soot LII signals in counterflow diffusion flames of 1-alkane fuels with 2-8 carbon atoms. Computed peak PAH concentrations were included. All data were normalized by the values for 1-butene flame. Reprinted from Wang et al. [635] with permission of Elsevier.

In addition to the above studies focusing on conventional hydrocarbon fuels, there are also CDF-based investigations on PAH/soot formation of oxygenated/bio-derived fuels. For instance, using a GC system with micro-probe sampling, Frassoldati et al. [601] quantified the mole fractions of stable intermediate species in both n-propanol and iso-propanol CDFs with $Y_{F,0} = 0.3$, $Y_{O,0} = 0.233$, and $K_E = 97.5\text{s}^{-1}$. They found that the iso-propanol flame had significantly higher level of C$_4$H$_6$ and C$_3$H$_6$, both of which have a pronounced tendency to form aromatic species—and thus—soot [174, 635]. Singh
et al. compared the soot [84] and PAH [441] formation tendencies of four butanol isomers in CDF with $X_{F,0} = 0.29$ and $X_{O,0} = 0.90$ (diluted by $N_2$) and observed that tert-butanol was the most sooting fuel, followed in decreasing order by n-butanol, iso-butanol, and sec-butanol. Although their results regarding the relative sooting tendency between n-butanol and iso-butanol (i.e., n-butanol is more sooting than iso-butanol) differed from previous studies in premixed flames [231] and yield soot index (YSI) rankings measured in coflow diffusion flames [653], the authors did observe that iso-butanol had a higher tendency than n-butanol to form small-sized PAHs, consistent with quantitative benzene measurements in coflow diffusion flames [646, 654]. Interestingly, LIF signals with longer detection wavelength (> 450 nm, typically believed to represent larger PAHs of more than four rings) showed that n-butanol has a greater tendency to form larger PAHs [441], in agreement with the ranking for sooting tendency. While no definite explanations were provided of this interesting trend reversal, it must be pointed out that in Ref. [441], due to large $X_{O,0}$ of 0.9, the flames from Singh and coworkers featured a rather different stoichiometric mixture fraction, and thus different flame structures, than the coflow fuel-air flames. It is likely that the ranking of sooting tendencies are dependent upon flame structures.

Sarathy and coworkers [589, 655] compared the chemical structures of methyl-butanoate (CH$_3$CH$_2$CH$_2$C(=O)OCH$_3$) and methyl-crotonate (CH$_3$CH=CHC(=O)OCH$_3$) diffusion flames in a counterflow configuration. The experimental results showed that the methyl-crotonate flame consistently produced higher levels of unsaturated species, such as C$_2$H$_2$, 1-C$_3$H$_4$, 1-C$_4$H$_8$, 1,3-C$_4$H$_6$, as well as C$_6$H$_6$, and is thus believed to have a higher propensity to soot than the methyl-butanoate flame. In a step further, Feng et al. [82] systematically measured the soot concentrations in CDFs for a class of model biodiesel fuels, emphasizing the effects of the type of ester functional group, number of carbon atoms, and double bonds. Since some of the fuels tested were of high molecular weight (up to methyl-undecanoate), and thus with rather low vapor pressure, these fuels were highly diluted with $N_2$ ($X_{F,0} \sim 0.1$) to maintain a moderate partial pressure. To promote soot formation for sensitive
measurements, pure oxygen was used as the oxidizer stream, which rendered the flames with SFO characteristics [73]. The study, summarized in Figure 22, showed that for the same functional ester group, the maximum soot volume fractions in the CDFs increased with the increase in fuel carbon number. Also, for esters with the same number of carbon atoms, methyl esters exhibited less sooting tendency than ethyl esters, and the unsaturated esters were significantly more sooting than their saturated counterparts. Nevertheless, the location of the double bond did not notably affect sooting tendencies.

Figure 22. Peak measured soot volume fractions of CDFs of various ester fuels: effects of length of carbon chain (a); comparison between methyl- and ethyl- esters (b); and effects of fuel unsaturation (c), adapted from Feng et al. [82] with permission of Elsevier.
5.1.2 Counterflow diffusion flames with fuel mixtures

It is noted that there are even more studies utilizing fuel mixtures than single-component fuels in CDF-based soot investigations on fuel effects. This popularity of the study of fuel mixing effects may be attributed to the following: 1) Conventional petroleum-based fuels and natural gases are always complex fuel mixtures [656], making understanding compositional effects on soot formation of practical importance; 2) The chemical interactions of various fuel molecules can sometimes shed light on important kinetic pathways for soot formation that are elusive in studies of single component fuels (a notable example is the synergistic effects of ethylene and propane mixtures on soot formation [528], detailed later); 3) The sooting tendencies of some fuels (i.e., high MW aromatic fuels) may be too high for accurate experimental determination of their sooting characteristics, and to study the relative effects in such cases it is usually beneficial to mix a small amount of the target fuel to a baseline flame.

a) Effects of CO/H$_2$ addition

Du et al. [568] investigated the effects of CO and H$_2$ addition in the fuel stream on soot inception rates in CDFs of C$_2$H$_4$, C$_3$H$_8$ and n-C$_4$H$_{10}$ fuels. Because the additives can passively affect soot formation by altering fuel concentration and flame temperature (in addition to directly influencing kinetic reactions), the authors conducted additional contrast tests by adding inert species with physical properties comparable to the target dopant. Of special note is that helium was used to contrast hydrogen for the isolation of the chemical effects of the latter. This choice was justified, since helium has a high diffusivity, similar to H$_2$, while it was shown that the preferential diffusion effects of these highly mobile species play important roles in strained flames [657-659]. The tendency for soot formation in the CDFs was quantified by the critical (minimum) local strain rate ($K_{cr}$, as measured by laser doppler velocimetry), at which light scattering signals from soot particles disappear (i.e., soot
inception is inhibited [551, 660]). Therefore, a higher $K_{cr}$ implies a stronger sooting tendency. The experimental results, summarized in Figure 23, showed that the addition of CO inhibited soot formation in C$_2$H$_4$ flames, while its addition in C$_3$H$_8$ and n-C$_4$H$_{10}$ flames had an interesting non-monotonic effect. Further analysis showed that CO exerted its soot-inhibitive effects in C$_2$H$_4$ flames mainly through fuel dilution. On the other hand, the initial enhancement of the sooting tendency with CO addition in C$_3$H$_8$ and n-C$_4$H$_{10}$ flames was thought to be caused by its chemical effects through reaction CO + OH = CO$_2$ + H, which may alter the concentrations of H and OH radicals that are important for soot formation and oxidation, respectively. Nevertheless, the authors did not provide direct evidence of this chemical effect and acknowledged that such reasoning could not explain the inhibiting behavior of CO in C$_2$H$_4$ flames [568]. Interestingly, in a subsequent study of coflow diffusion ethylene flames by Guo et al.[387], a detailed kinetic model suggested that in addition to the dilution effect, CO plays a chemically-enhancing effect on soot formation by: 1) raising flame temperature; 2) increasing the surface growth rates due to higher H radical production; and 3) decreasing soot oxidation due to higher OH radical consumption. They noticed that the overall soot production rate was reduced, implying that the dilution effect was still dominant. However, no further study was conducted to explain the experimental results of Du et al. [568] that soot inception was enhanced by CO addition in C$_3$H$_8$ and n-C$_4$H$_{10}$ CFD flames. CO may have a role in influencing fuel pyrolysis chemistry, but further studies are needed.
Figure 23. Effects of CO/H$_2$ addition in the fuel side on critical strain rates for soot formation in CDFs with base fuels of C$_2$H$_4$ (a); C$_3$H$_8$ (b) and n-C$_4$H$_{10}$ (c). Adapted from Du et al. [568] with permission of Elsevier.

The addition of H$_2$ was found to decrease the sooting tendencies of CDFs of C$_2$H$_4$, C$_3$H$_8$ and n-C$_4$H$_{10}$ fuels [568], even though flame temperatures were always enhanced. The following were believed to offset soot-promoting effects of higher flame temperature from H$_2$ addition: 1) In positively stretched flames, the preferential diffusion effects of H$_2$ may cause precursor concentration reduction, decreasing the rate of soot formation; 2) In the HACA framework for PAH growth [661], the addition of H$_2$ may push the hydrogen abstraction reaction A$_i$ + H = A$_i$- + H$_2$ backward, preventing
aromatic radicals (A\textsubscript{i-}) from participating in further mass addition reactions, and leading finally to reduced soot precursor concentrations; 3) The addition of H\textsubscript{2} also decreases the fuel carbon supply and thus effectively plays a role of dilution. As previously mentioned, comparison against helium addition cases may help isolate the above effects. However, it has been noted that the relative effectiveness of H\textsubscript{2} and helium in inhibiting soot formation is not always consistent. For example, in the CDFs of Du \textit{et al.} [568], helium was more effective than hydrogen in suppressing soot inception for all the three fuels tested, consistent with a study on methane coflow flames [662]. But studies in ethylene/air coflow flames [663, 664] showed exactly the opposite, i.e., hydrogen was more effective than helium in reducing soot formation. Compared to helium, the addition of hydrogen increases flame temperature while it may also inhibit PAH/soot HACA growth. Thus, it is highly likely that the dominating effect is dependent upon individual flame conditions. Obviously, more studies are needed to clarify these interesting phenomena.

Through visual observation of soot luminosity, Choi \textit{et al.} [624] also noticed a reduction of soot formation by H\textsubscript{2} addition in ethylene-air CDFs. Thermophoretic sampling of the soot particles and subsequent analysis with HRTEM showed that the size of the particles was reduced, while the formation of fullerenic nanostructures (represented by the curved carbon lamella) was enhanced by the addition of hydrogen. However, the exact location where soot extraction was performed in the flames was not reported. The authors postulated that the enhanced curved structure may have been a result of enhanced five-member ring formation after hydrogen addition, although no further evidence was shown.

\textbf{b) Synergistic effects of ethylene-based binary fuel mixtures}

The influence of fuel-stream mixing of ethylene and propane on soot formation was investigated by Hwang \textit{et al.} [528] in CDFs with \(X_{\text{F,0}} = 1.0, X_{\text{O,0}} = 0.24\) and \(K_E = 55\text{s}^{-1}\) at various fuel mixing ratios. Soot volume fractions and relative PAH concentrations were measured with light extinction/scattering and laser-induced fluorescence, respectively. An interesting synergistic effect
was observed, i.e., PAH/soot formation was enhanced in the flames of ethylene/propane mixtures compared to both neat ethylene and neat propane flames. Specifically, the PAH/soot concentrations of the ethylene-propane flames were observed to increase first and then decrease with more propane mixing, as can be seen in Figure 24. It was argued that the monotonic changes in flame temperature and C₂H₂ concentrations with the propane mixing ratio cannot explain such non-monotonic variation in PAH/soot formation, so that chemical cross-linking between the fuel molecules must play a role. In this regard, kinetic modelling of the flames was conducted and the results showed that the peak concentration of C₃H₃ (known to be an important benzene precursor [152]) also exhibited a non-monotonic variation with the propane mixing ratio. Further kinetic analysis [528, 616] revealed that the relatively low concentration of C₃H₃ in the sooting region of the neat ethylene flame acted as a bottleneck to the formation of aromatic soot precursors, while with a small amount of propane addition, the concentrations of methyl radicals (CH₃) increased as a direct consequence of propane decomposition. The increased CH₃ concentrations then promoted various reactions between C₁ and C₂ species, leading to an enhancement of C₃H₃ [642], and finally--benzene production. As an important bridge toward large PAHs and soot particles, the increase in benzene concentration translated to the observed increase in PAH/soot when a small amount of propane was added to the ethylene baseline flame. With further increases in the propane mixing ratio, the concentration of C₂H₂ decreased notably, contributing negatively to PAH/soot growth through the HACA mechanism and leading to a decrease in sooting tendency. In this way, the study by Hwang et al. [528] demonstrates the interplay between C₃H₃ and C₂H₂ in determining the final soot loading of ethylene-propane CDFs, with C₃H₃ controlling incipient aromatic ring formation and C₂H₂ dominating large PAH and soot mass growth. In a subsequent effort to identify the role of C₃ chemistry in PAH growth beyond benzene, Lee et al. [582] measured PAH and soot concentrations in CDFs of ethylene-propane mixtures with a small amount of benzene addition (1.83%, molar concentration) to the fuel stream. It was argued that benzene doping would significantly increase benzene concentration in the pyrolysis...
and flame zone, thereby mitigating the relative importance of the effect of C$_3$ pathways on initial ring formation to the ethylene/propane synergistic behaviors. LIF signals, detected at different wavelengths, showed that although the synergistic effect nearly disappeared in small aromatic species (represented by LIF signal with 330 nm detection wavelength), it was still pronounced in relatively larger PAHs (>400 nm detection wavelength). These results suggested that adding propane to ethylene affects not only the incipient ring formation but also PAH growth beyond benzene.

![Figure 24](image_url)

**Figure 24.** Synergistic effect on PAH and soot in ethylene-propane flames. Shown are peak soot volume fraction and PAH LIF signals as a function of propane mixing ratio. A non-monotonic variation can be clearly identified. Reprinted from Hwang *et al.* [528] with permission of Elsevier.

In addition to the influence on benzene formation through C$_3$H$_3$ recombination, Trottier *et al.* [665] argued (based on combined experimental and numerical investigations on soot formation in coflow diffusion flames of binary fuel mixtures) that the increase in CH$_3$ radical from propane in ethylene-propane flame may enhance the forward reaction of C$_2$H$_4$ + CH$_3$ = C$_2$H$_5$ + CH$_4$, increasing the concentration of vinyl radicals (C$_3$H$_3$), which in turn contribute to higher benzene production through various C$_4$ + C$_2$ channels. This argument was supported by numerical analyses of Liu *et al.* [284], Mc Nesby *et al.* [101], and Park *et al.* [666].
It is important to note that for both the $C_3 + C_3$ or $C_2 + C_4$ benzene formation pathways, the CH$_3$ radical, a direct pyrolysis product from propane decomposition, always plays a critical role in explaining the synergistic effect seen in ethylene-propane mixtures. It can thus be reasoned that the sooting tendency of ethylene flame may also be enhanced by other fuel dopants, as long as such dopant can decompose to produce appreciable amounts of CH$_3$ radicals in the fuel pyrolysis zone. If—as a neat fuel—the dopant also has a lower sooting tendency than ethylene, synergistic effects are then expected to develop between ethylene and this dopant. Besides C$_3$H$_8$, many other fuels, including CH$_4$, C$_2$H$_6$, n-butane, iso-butane, dimethyl ether (CH$_3$-O-CH$_3$, DME), ethanol (C$_2$H$_5$-OH) and n-propanol are known to meet these criteria. And indeed, the synergistic effects on PAH/soot formation have been observed for C$_2$H$_4$/C$_2$H$_6$ [586], C$_2$H$_4$/n-C$_4$H$_{10}$ [86], C$_2$H$_4$/i-C$_4$H$_{10}$ [86], C$_2$H$_4$/DME [90, 100, 476], C$_2$H$_4$/C$_2$H$_5$OH [41, 101, 607, 667], C$_2$H$_4$/C$_3$H$_7$OH [666] mixtures, as summarized in Figure 25. The peak soot volume fractions in CDFs of these ethylene-based binary mixture fuels were plotted as a function of molar mixing ratio, after being normalized by their respective values in the neat ethylene flames. Although not all the above studies were accompanied by detailed kinetic investigations, the analyses of reaction pathways for those with unanimously pointed to the role of CH$_3$ in inducing synergistic effects (through C$_1$ and C$_2$ interactions, as discussed previously). The strength of the synergistic effects (i.e., the quantitative extent by which the dopant can increase PAH/soot formation as compared to the ethylene baseline flame) were seen to be largely dependent on the relative ease for the dopant to decompose to CH$_3$ radicals in the flame pyrolysis zone. For instance, the fact that 5% of propane addition in ethylene CDF increased the peak SVF by 32% (larger than the 14% as induced by 5% ethane addition) can be partially explained by the higher CH$_3$ concentration in the case of propane doping [140]. The stronger synergistic effect on benzene and soot formation of ethylene/DME mixture than in ethylene/ethanol mixture flames can also be rationalized by the higher production rates of CH$_3$ in DME pyrolysis than ethanol [642].
The effect of CH₄ addition in C₂H₄ CDF deserves special attention. The sooting tendency of CH₄ is known to be the lowest among all hydrocarbon fuels, which may be rationalized by the fact that CH₄ has the highest H/C ratio and there is no C-C bond in the CH₄ molecule. Nevertheless, considering that CH₃ can be formed by hydrogen abstraction from CH₄, it is expected that a small amount of CH₄ addition in C₂H₄ flame would enhance soot formation. However, a relatively monotonic decrease in soot volume fraction with CH₄ mixing ratio was observed experimentally in ethylene CDFs [586]. On the contrary, both experiments [586] and numerical simulation [140] suggested increased PAH formation by CH₄ addition, although its quantitative extent was rather small compared to cases with other dopants (e.g., C₂H₆ and C₃H₈). Yoon et al. [586] explained this interesting behavior of CH₄ doping based on the following: First, the C-H bond energy (483.6 kJ/mol) in CH₄ is notably higher than the C-C bond energy in C₂H₆ (376.1 kJ/mol) and C₃H₈ (371.3 kJ/mol), so it is more difficult for CH₄ to break into CH₃ radicals than for C₂H₆ and C₃H₈, especially in the low temperature region (1000–1500K, fuel side of the flame) where PAHs were formed. Therefore, the concentration of CH₃ in the methane mixing case was rather low, resulting in the weaker synergistic effect on PAH formation compared to other cases. Second, the flame with CH₄ mixing
has lower flame temperature and C\textsubscript{2}H\textsubscript{2} concentrations, which in turn lead to a lower soot surface HACA growth rate than in the C\textsubscript{2}H\textsubscript{6} and C\textsubscript{3}H\textsubscript{8} mixing cases. The slight increase in PAH concentration with CH\textsubscript{4} mixing could not compensate for this reduction in surface growth rate and, finally, a decrease in soot loading resulted. It is worthwhile to contrast this monotonic decrease of SVF in CDFs to an earlier investigation using coflow experiments, where a notable synergistic effect of ethylene and methane mixture was observed on peak centerline soot volume fraction [668]. Although the difference may have been caused by the different time-temperature histories encountered between the counterflow and coflow configurations, it was noted that in the coflow study [668] the fuel was progressively less diluted with N\textsubscript{2} from the neat C\textsubscript{2}H\textsubscript{4} to the neat CH\textsubscript{4} case, so that the different level of dilution may also play a role. More studies are required to unravel this discrepancy.

The special role of C\textsubscript{2}H\textsubscript{4} was rather evident, as it was involved in all the above cases that exhibited synergistic effects in binary fuel mixtures. McEnally and Pfefferle [642] insightfully noted that since it is not easy for C\textsubscript{2}H\textsubscript{4} to decompose to CH\textsubscript{3}, the sooting characteristics of C\textsubscript{2}H\textsubscript{4} flame are particularly sensitive to CH\textsubscript{3} produced from the dopants. As a matter of fact, C\textsubscript{2}H\textsubscript{4} (along with C\textsubscript{2}H\textsubscript{2}) is special, since most other common alkane and alkene fuels can easily form CH\textsubscript{3} in the fuel pyrolysis process. It can thus be reasoned that no CH\textsubscript{3}-induced synergistic effect may be developed for flames of base fuels other than C\textsubscript{2}H\textsubscript{4} and C\textsubscript{2}H\textsubscript{2}. Indeed, with an increasing amount of DME (CH\textsubscript{3}-O-CH\textsubscript{3}, an effective source of CH\textsubscript{3} radical) was added to CDFs of CH\textsubscript{4}, C\textsubscript{2}H\textsubscript{6} and C\textsubscript{3}H\textsubscript{8}, the peak soot volume fraction and PAH LIF signals showed only a trend of monotonic decrease [100], which can be understood since all of the above base fuels can–by themselves–produce appreciable amounts of CH\textsubscript{3} and thus are not sensitive to additional CH\textsubscript{3} from DME decomposition. One notable exception that cannot be explained by this reasoning is the synergistic effects of methane and ethane mixtures, as observed in coflow diffusion flames of Trottier \textit{et al.} [665]. However, considering that no quantitative soot volume fraction data or kinetic explanation was provided in Ref. [665] with respect to this fact, it
may be interesting to perform additional experimental and kinetic modelling studies to confirm such a trend in CDFs.

As demonstrated above, non-monotonic variation of PAH and soot concentrations with mixing ratio have been widely observed in both counterflow and coflow [71, 284] diffusion flames of ethylene-based binary fuel mixtures. However, the situation in premixed flames can be drastically different. For instance, Wu et al. [103] measured PAH and soot in rich premixed flames of ethylene-ethanol mixtures and observed that ethanol addition monotonically decreased soot formation. Similar observations were made by Salamanca et al. [607] for ethylene-ethanol, and by Sirignano et al. [90] for ethylene-DME premixed flames. These interesting results suggest that physicochemical pathways of PAH/soot formation for the same fuels can differ notably, depending on flame conditions. A hint of such dependence (in particular, the presence (absence) of the synergistic effects of ethylene-based binary fuels in non-premixed (premixed) flames) may be found in the results from partially-premixed CDFs. Hwang et al. [165] found that the non-monotonic variation of soot volume fraction in CDFs of ethylene-propane mixtures can be mitigated–and even eliminated–by premixing the fuel stream with oxygen. The explanation lay in the fact that with an oxidizer present in the fuel stream, the oxygen atom may react in the fuel pyrolysis zone with C₂H₂ to form CH₂ radical (C₂H₂ + O = CH₂ + CO), which in turn interacts with C₂H₂ to form the important benzene precursor, propargyl radical (C₂H₂ + CH₂ = C₃H₃ + H). In such a way, the bottleneck of benzene formation, caused by the deficiency of C₁ species in neat C₂H₄ flame, can be effectively removed and as a result, the partially-premixed C₂H₄ flame would not be as sensitive to additional CH₃ from propane decomposition, mitigating the synergistic effect. Although yet to be confirmed, the monotonic decrease in soot concentration in ethylene-ethanol [103] and ethylene-DME premixed flames [90] is thought to be explainable by this mechanism.
c) Surrogates for practical fuels

Practical fuels are complex mixtures of hundreds of hydrocarbons [656, 669-672] that prohibit kinetic simulation with detailed chemistry models. Therefore, simplified surrogates with a relatively small number of components are needed to represent various physical and chemical properties of real fuels. As a critical fuel property, the sooting tendency is an important consideration in surrogate development [673, 674], rationalizing the wide interest in the compositional effects of surrogate fuels on soot formation. Often the studies of surrogate fuels were performed by doping a small amount of the surrogate into the baseline flames of simple hydrocarbons (i.e., methane or ethylene), so that the compositional effect could be better isolated from other non-chemical factors, such as flame temperature and flow fields [123, 675, 676]. In addition, the risk of fuel vapor re-condensation can be minimized since the partial pressure of a target fuel may be rather small [677]. This point is particularly important for fuel components with a high molecular weight, and thus, a high boiling temperature.

With in-flame gas sampling and GC/MS analysis, Bufferand et al. [677] compared the detailed chemical structures of methane counterflow diffusion flames doped with 1000 ppm of real jet fuel and a corresponding six-component surrogate (Utah/Yale surrogate). With highly-diluted non-sooting flames to facilitate probe sampling, the experimental data showed that the jet fuel-doped flame produced notably more benzene and toluene (more than a factor of two) than the surrogate-doped flame, although the surrogate was seen to capture other general jet fuel behavior reasonably well. In a follow-up work, Jahangirian et al. [80] investigated incipiently sooting ethylene counterflow diffusion flames doped with 2000 ppm of jet fuel and two surrogates (six-component Utah/Yale surrogate [677] and two-component Aachen surrogate). Good agreement was observed between the real fuel and the surrogates in terms of peak location and magnitude of benzene concentration. With respect to toluene formation, the Utah/Yale surrogate performed better in mimicking the behavior of
real jet fuel. Nevertheless, it was concluded that both surrogates exhibited acceptable agreement with real fuel in terms of aromatic formation under incipient sooting conditions.

Realizing the lack of fundamental studies in non-premixed flames of critical transport/surrogate fuel components, Carbone and Gomez conducted a series of investigations on the chemical structures of CDFs doped with hundreds of ppm of toluene [606], 1,2,4-trimethyl benzene (TMB) [609], n-decane [617] and n-decane / TMB mixtures [621]. In these studies, two baseline flames (i.e., a non-sooting methane SFO flame and an incipiently sooting ethylene SF flame) with significantly different thermal-chemical environments were used as flame reactors to provide well-defined (and contrasting) environments for the examination of reaction pathways of the target fuels. Profiles of flame temperature and mole fractions of important intermediate species were measured with thermocouples and GC/MS techniques, respectively. Analysis of the experimental data, together with numerical results obtained with literature-based kinetic mechanisms, also provided direction for future improvement of chemical models, especially in the application of non-premixed flame coupling kinetics and transport.

Choi et al. [622] doped n-heptane, benzene, and toluene separately, up to 20% (molar ratio) in sooting ethylene CDF \( X_{F,0} = 1.0, X_{O,0} = 0.3 \), and quantified the effects of these dopants on PAH and soot formation with the techniques of LIF and LII, respectively. The addition of benzene and toluene was confirmed to enhance soot/PAH formation, with toluene exhibiting stronger effects, which was consistent with the higher sooting tendency of toluene [678]. Kinetic analyses revealed that in toluene-doped flames, the formation pathway for large PAHs from benzyl radicals through indenyl radical \( A1CH_3 \rightarrow A1CH_2 \rightarrow C_9H_8 \rightarrow C_9H_7 \rightarrow A4 \) played a significant role, and was believed to partially explain the higher soot loading of toluene.

With similar measurement techniques, the sooting characteristics of n-heptane, iso-octane and toluene, components of the widely-used toluene primary reference fuel (TPRF), were studied in CDFs by Choi et al. [447]. Both SF and SFO flames were tested and the liquid fuels were diluted only by
N$_2$, without employing a base gaseous fuel. As can be seen in Figure 26, the fuel and oxidizer dilution levels for the SF flames were somewhat high ($X_{F,0} = 0.30$ and $X_{O,0} = 0.21$), so the flame was non-sooting when neat n-heptane or iso-octane was the fuel. Soot loading was monotonically increased as more toluene was added, and the behavior was qualitatively similar between n-heptane/toluene and iso-octane/toluene mixtures. Note, however, that the effect of toluene addition only became significant when the toluene ratio ($R_T$) was larger than 0.4. On the other hand, interesting non-monotonic variations of PAH concentrations (represented by the intensities of PAH LIF signals) with $R_T$ were observed in both SF and SFO flames, which were qualitatively captured by kinetic modelling using a mechanism based on the literature [679]. It was determined through analysis of the numerical results that, as $R_T$ increased, the concentration of benzyl radical increased, while that of the H radical decreased, leading to non-monotonic variations of the rates for major benzene production reaction: $A1CH_3 + H = A1 + CH_3$. In addition, ethynylbenzene ($A1C_2H$) showed significant synergistic effects, while biphenyl radical (P2-) increased and $C_2H_2$ decreased with an increase in $R_T$, resulting in the non-monotonic behavior of $A3$ through $A1 - A1C_2H = A3 + H$ and $P2 - C_2H_2 = A3 + H$, respectively. While this investigation [447] dealt with only binary mixtures, a follow-up study extended the work to ternary mixtures [629].

![Figure 26](image.png)

**Figure 26.** Normalized peak soot volume fraction (measured with LII) and PAH LIF signal as a function of toluene mixture ratio ($R_T$) in counterflow diffusion flames of n-heptane/iso-octane and
toluene mixtures. Fuel and oxidizer are both diluted with N\textsubscript{2} and \( X_{F,0} = 0.30, X_{O,0} = 0.2 \). Reprinted from Choi et al. [447] with permission of Elsevier.

5.1.3 Quantitative indices of sooting tendencies

The above discussions demonstrate that fuel molecular structures are critical in determining soot production. This has important implications for the prediction of soot emissions from practical combustion devices which typically burn real fuels (i.e., diesel and jet fuels) with very complex compositions [669, 670]. It is common for current predictive tools to use surrogates to represent real fuels [669, 670, 673], making a quantitative index necessary to assess whether the sooting tendencies among them are matched [680]. A sooting index can also be helpful for comparing sooting tendencies among fuels with different functional groups, so that statistical information on the relation between fuel molecular structure and sooting tendency can be obtained [678, 681, 682]. This sub-section briefly reviews available sooting indices, with particular focus on the sooting temperature index (STI), which is based on sooting limits of counterflow diffusion flames.

Sooting metrics have been proposed for pyrolysis, premixed, and non-premixed combustion conditions. Crossley et al. [683] pyrolyzed 20 μL of vaporized liquid samples in an alumina bed at 850ºC and quantified the amount of carbon deposition with a temperature-programmed oxidation system. By normalizing the soot deposition of different fuel samples to two reference compounds, a micropyrolysis index (MPI) was proposed, which was believed to provide information on the tendency of a particular fuel to soot through oxygen-free pyrolysis (i.e., independent of heating value, oxidation chemistry, oxidizing environment, and etc.).

For premixed systems, critical equivalence ratios, or C/O ratios at which soot luminosity can be visually detected, were frequently used as a quantitative metric [119, 645, 684]. Considering that under fuel-rich conditions, fuel carbon is converted primarily to CO instead of CO\textsubscript{2}, a critical effective equivalence ratio was also proposed [685], based on the stoichiometric relation in which CO and H\textsubscript{2}O were the combustion products.
Sooting tendencies in non-premixed systems have been historically quantified using smoke point, which is the height of the highest flame without soot leakage when the test fuel is burned in a wick-fed lamp [686], or in a coflow burner (for gaseous fuels) [104, 117]. Laminar co-flow jet flames are characterized by a buoyancy-controlled velocity field so that the velocity at a given height above the burner is proportional to the square root of the height, while reasonably independent of the fuel outlet velocity or volumetric flow rate [66]. As a result, as the fuel flow rate (or outlet velocity) increases, the flame becomes longer, increasing the fuel residence time, which in turn enhances the sooting propensity of the flame. There exists a critical flame height above which soot formed in the fuel-rich region of the flame cannot be fully oxidized and smoke begins to pierce from the flame front to emit into the environment. This critical flame height is typically defined as the smoke point. Obviously, fuels with a lower smoke point have a higher tendency to soot and there is a historical sooting index that was defined to be inversely proportional to the smoke point [687].

Quantitative determination of the critical flame height (i.e., the smoke point) is somewhat subjective and can vary from one instrument to another. Furthermore, as more oxygen is needed to form a stoichiometric fuel/oxidizer condition at the flame front higher molecular weight fuels naturally have longer flame [680], irrespective their sooting tendencies. In this regard, smoke point is affected by both the sooting tendency and the molecular weight of the fuel. In an effort to reduce this undesirable molecular weight effect, Calcote and Manos [645] introduced the threshold soot index (TSI), with a scale of 0 (ethane) to 100 (naphthalene), where higher values mean higher sooting propensity. This index is a normalization of smoke point data as given by:

\[
TSI = a \left( \frac{MW}{SP} \right) + b
\]

Eq. 24

where MW is the molecular weight of the test fuel, SP is the smoke point and \(a\) and \(b\) are apparatus-specific constants. Note that by considering the linear relationship between flame height and fuel volumetric flow rate, \(SP\) can also be replaced by the critical fuel volumetric flow rate in Eq. 24 for the definition of TSI. Despite its wide use, TSI is just one way among many to normalize smoke point
data. Arguing that the direct inclusion of MW in sooting index definition is not physically motivated, Li and Sunderland [688] revisited smoke point data and proposed a new sooting metric called the normalized smoke point (NSP). Barrientos et al. [681] proposed the oxygen extended sooting index (OESI) for oxygenated fuels to account for the effect of oxygen atoms in the normalization of smoke point data.

Another sooting index that received widespread attention is the yield sooting index (YSI), as pioneered by McEnally and Pfefferle [689]. They doped target fuels in a methane coflow diffusion flame and measured the peak soot volume fraction, which was then normalized to determine YSI [67, 653, 678, 690, 691]. YSI was traditionally obtained by measuring SVF using the LII technique, and a more recent study [692] has shown that the techniques of color-ratio pyrometry and spectral soot emission are also compatible. In addition, measured YSI values were also unvarying across flame conditions with different temperatures and levels of partial premixing, suggesting that the robustness of YSI to be used across a range of premixed and temperature conditions is relevant to soot formation [692].

The above indices (except the MPI) are based on either coflow or smoke point flames in which undiluted fuel and air were typically used. Recent research in novel combustion techniques such as oxygen-enriched combustion, low temperature combustion and flameless combustion [693] necessitates the study of the effects of fuel and oxidizer stream dilution on soot characteristics of diffusion flames. Furthermore, considering soot formation is a rate-limiting process [554], and therefore sensitive to the characteristic flow time scale, it would be ideal if the effects of residence time are reflected in the sooting index. In coflow diffusion flames, the residence time is typically too long to represent combustion under practical conditions [114]; challenges also exist for its active control. In addition, as discussed in a previous section, in coflow flames, soot once formed will be convected toward the oxidizing zone where soot oxidation takes place, behaving as a soot formation/oxidation (SFO) flame. In such a case, the smoke point is essentially a critical condition
when soot formation and oxidation are balanced. However, to quantify the sooting tendency for various fuels, it may be helpful to avoid the interference from soot oxidation.

CDF is a flame configuration that can provide valuable information regarding residence time effects on soot formation; because it is relatively easy to parameterize the strain rate (inversely proportional to residence time) by adjusting the nozzle exit velocities. Also, by utilizing SF flames, it is possible to separate the soot formation process from oxidation. The sooting tendencies of different fuels have been systematically studied in CDFs in the form of sooting limit curves [620]. An example sooting limit map is shown in Figure 27 for various C_1-C_4 fuels. A CDF can be manipulated to be sooting or non-sooting by varying the amount of N_2 dilution in the fuel or oxidizer stream (detailed discussion on dilution effect on soot formation are presented in Section 5.2).

Experimentally, whether a CDF is sooting can be determined by comparing the Rayleigh and Mie scattering signals from gas molecules and soot particles, respectively. As detailed in Ref. [614], under non-sooting conditions the Raleigh scattering intensity varies monotonically from the fuel to the oxidizer stream due to the change in the scattering cross-section of gases. Once soot is formed, the Mie scattering signal from soot is embedded in the Rayleigh signal, resulting in a local maximum. The presence of this local maximum is indicative of the onset of soot.

It was shown in Ref. [614] that in an ethylene CDF with X_{F,0} = 1.0, soot formation could only be observed when X_{O,0} was higher than 0.18. As such, X_{F,0} = 1.0 and X_{O,0} = 0.18 can be regarded as a critical condition for soot formation. Similarly, a corresponding critical X_{O,0} always exists for a specified X_{F,0}. In this regard, a sooting limit curve can be drawn in the X_{F,0}–X_{O,0} plane by connecting the critical sooting limit points (X_{F,cr}, X_{O,cr}) as shown in Figure 27.
Figure 27. Sooting limit map of various C₁-C₄ fuels in counterflow diffusion flames, measured with nozzle exit velocity at 40 cm/s. Reprinted from Wang and Chung [620] with permission of Elsevier. Brown dotted line represents theoretical SF–SFO flame boundaries.

These sooting limit curves divide the $X_{F,0} - X_{O,0}$ plane into two regions: The upper right is the sooting region, and the remainder is the non-sooting region. Any combination of $X_{O,0}$ and $X_{F,0}$ in the upper right region will result in a sooting flame. It is clear from Figure 27 that the sooting regions have different sizes for different fuels. For example, the sooting region for C₂-C₃ alkenes are noticeably larger than their alkane counterparts and iso-butane has a more extended sooting zone than n-butane. These are consistent with the known fact that, in diffusion flames, alkenes are generally more sooting than alkanes [99], and branched alkanes are more sooting than those with straight chains (with the same carbon number) [688]. A more interesting observation can be made when re-plotting the sooting limit data shown in Figure 27 as a function of calculated flame temperature $T_f$. The results are shown in Figure 28. It can be seen that for SF flames (i.e., large $X_F$), the flame temperature remains nearly fixed over a wide range of $X_{F,cr}$, indicating the existence of a threshold temperature for the onset of soot formation in SF flames. The flame will be sooting if it has a peak temperature above this threshold, and vice versa. What is more important is that the threshold temperature is highly dependent on the fuel type, with less sooting fuel featuring a higher threshold temperature at the sooting limit.
Figure 28. Critical fuel mole fractions at sooting limit conditions plotted as a function of calculated flame temperature for various C$_1$-C$_4$ fuels in counterflow diffusion flames with nozzle exit velocity at 40 cm/s. Reprinted from Wang and Chung [620] with permission of Elsevier.

In fact, this dependence of the threshold temperature for SF flame with $X_{F,0} = 1.0$ ($T_{cr;SF}$) on fuel type provides an alternative means for ranking the sooting tendency of different fuels. To demonstrate and validate the possibility for its use as a quantitative sooting index, the correlation between $T_{cr;SF}$ and the threshold soot index (TSI) is presented in Figure 29, where good correlation can be seen. This suggests that $T_{cr;SF}$ determined from counterflow experiments can be a viable metric for sooting propensity. Note that the uncertainties of determining $T_{cr;SF}$ are considerably lower than that for TSI.
Figure 29. Correlation between threshold temperature for soot formation in SF flames and literature data of TSI. Reprinted from Wang and Chung [620] with permission of Elsevier.

By normalizing $T_{cr;SF}$ of the test fuels to two reference fuels in a similar manner as used for YSI, a sooting temperature index (STI) can be defined. This CDF-based sooting index has the following novel features compared to smoke point-based ones: 1) It quantitatively represents the limiting conditions of soot onset for a target fuel, while smoke point represents a condition where soot loading is too excessive to be fully oxidized; 2) In the test environment for SF flames, interference from soot oxidation is absent, so the sooting tendency can be assessed purely on the fuel’s chemical reactivity to form soot; 3) The effect of residence time can easily be incorporated quantitatively in STI, by measuring sooting limits with different nozzle exit velocities. A sooting sensitivity index (SSI) was also proposed by normalizing the slope of $T_{cr;SF}$ as a function of nozzle exit velocity ($dT_{cr;SF} / dV_0$), and it was found that the sooting tendency of different fuels also has different sensitivity to variations in residence time [620].

Finally, it may also be noteworthy that the approach used to develop YSI [689] (i.e., doping a small amount of test fuel in a base flame and measure the peak SVF) can also be applied to CDF, and further efforts are needed to quantify the dependence of sooting tendency data especially on pressure.
5.2 Effects of dilution and non-hydrocarbon additives on soot formation

The identification of soot-inhibiting non-fuel additives is an important research area for practical purposes [564]. For example, flue gas recirculation (FGR), which is essentially the addition of CO₂ and H₂O in reactants, is frequently used in utility boilers as a measure for emission reduction [694]. Similarly, exhaust gas recirculation (EGR) is widely adopted in internal combustion engines. Oxy-fuel combustion is also actively explored to reduce the cost of CO₂ sequestration. It is of fundamental interests to understand the underlying mechanisms on how these additives affect soot formation. Typical non-hydrocarbon additives include inert gases such as nitrogen, argon, helium, carbon dioxide, and water vapor, as well as oxidizing agents of oxygen (i.e., partial-premixing), ozone, and nitrogen dioxide. The influences of organometallic and halogenated compounds on soot formation have also attracted research interests. In this section, relevant studies conducted with laminar counterflow flames are reviewed and discussed.

5.2.1 Nitrogen and noble gas addition

The effect of nitrogen dilution in oxidizers was studied by Vandsburger et al. [94] in counterflow diffusion flames of ethylene and propane, stabilized in the forward stagnation region of a cylindrical porous burner. The level of oxidizer dilution was quantified by \( X_{O,0} \). The flames were comprehensively characterized; the profiles of axial velocity (LDV), temperature (thermocouples), and soot volume fraction, number density, and average diameter (light extinction and scattering) were all measured along the stagnation streamline. The results showed that for ethylene flames, the increase in \( X_{O,0} \) from 0.18 to 0.28 resulted in increases in the maximum flame temperature (1750 K to 2150 K) and peak soot volume fractions (from \( 3.7 \times 10^{-7} \) to \( 2.1 \times 10^{-6} \)). Maximum particle number densities (maximum particle sizes) were measured to occur close to the flame front (stagnation plane), and increased slightly with \( X_{O,0} \). Propane flames exhibited similar behavior with respect to the effect of \( X_{O,0} \), although soot loading was notably lower compared to ethylene. Hwang and Chung [281]
performed similar experiments in ethylene SF CDFs and found that both soot and PAH formation were enhanced with the increase in $X_{O,0}$. In a more recent work [614], the range of $X_{O,0}$ investigated was extended further, up to 0.5 for ethylene CDF ($X_{F,0}=1$), and similarly, a monotonic increase in soot loading with increased $X_{O,0}$ was observed. In a counterflow diffusion flame with neat methane as the fuel stream, Beltrame et al. [576] increased $X_{O,0}$ from 0.21 all the way to 1.0 (pure oxygen). Their experimental and numerical results agreed, showing a monotonic increase in peak soot volume fraction. As a related fact, it is interesting that Kennedy and coworkers also observed the formation of multi-walled carbon nanotubes [506, 695], as well as fullerenes [696] in methane or acetylene CDFs with oxygen-enriched oxidizer.

At first glance, it is intriguing that an increase in oxygen content in the oxidizer stream (i.e., $X_{O,0}$) would strengthen soot formation. It might seem that the increased amount of oxygen would enhance soot oxidation, and therefore, result in lower soot production. This can be true for over-ventilated coflow diffusion flames, where the final soot loading is determined by competing effects of soot production in the fuel-pyrolysis zone, and subsequent soot oxidation in the oxygen-rich post-flame zone. For example, Glassman and Yaccarino [697] found that the sooting tendency of ethylene coflow diffusion flame (represented by the normalized sooting flame height) decreased as $X_{O,0}$ increased from 0.16 to 0.24, which was explained by the fact that higher $X_{O,0}$ resulted in a higher soot burn-up rate. However, all the counterflow flames mentioned above are of the SF type, so that the flame was stabilized on the oxidizer side of the stagnation plane. As discussed in a previous section, in such cases the soot particles would be transported away from the oxidizing flame sheet so that soot oxidation was largely absent. Higher $X_{O,0}$ would lead to higher temperatures in the reacting flame front as well as in the fuel pyrolysis zone, enhancing fuel pyrolysis and soot formation rates [93, 698].

The effect of fuel stream dilution by inert gases has also been studied. Du et al. [568] measured the critical strain rate for soot inception (i.e., soot inception limit, $K_{cr}$) in CDFs of C$_2$H$_4$, C$_3$H$_8$, and n-C$_4$H$_{10}$ with fuel side dilution by nitrogen and argon. The data for the C$_2$H$_4$ flame is re-plotted in Figure
30, and other flames exhibit similar qualitative trends. As can be seen, and much as expected, $K_{cr}$ decreased (i.e., sooting tendency decreased) monotonically with the increase in the dilution level. Considering that inert dilution can affect soot formation by variation of either the flame temperature or the concentration of soot precursors, additional efforts were taken to isolate these effects. In Figure 30b, the dashed line represents the variation of $K_{cr}$ with flame temperature, which was varied by substituting part of the nitrogen in the oxidizer with equal moles of argon, without altering the fuel/oxidizer dilution level [699]. In such a way, this line depicts only the thermal effect. The symbols are data shown in Figure 30a, re-plotted as a function of temperature. Its deviation from the dotted line indicates the inert affecting soot inception via not only thermal, but dilution effects. Indeed, the decrease in $K_{cr}$ with N$_2$/Ar addition was faster than expected from only thermal considerations. At a specified temperature, $K_{cr}$ (in cases with argon dilution), was even lower than those with N$_2$ dilution. This can be explained by the fact that more argon is required than nitrogen for the temperature to be matched, due to the lower specific heat of argon, so the argon diluted flame has a higher level of dilution. The importance of concentration modification in explaining the reduction of soot by inert addition was also noted in experiments of jet flames [96, 700].
Figure 30. Critical strain rates for soot formation in ethylene CDFs with fuel side N₂ and Ar dilution (a) and effects of temperature on soot inception (b). Replotted with data taken from Du et al. [568].

In addition to $K_{cr}$, sooting characteristics were also compared carefully in moderately sooting CDFs, with and without fuel stream inert addition [74]. Soot volume fraction, axial velocity and temperature profiles were experimentally determined for ethylene CDFs with $X_{F,0} = 1.0$, 0.75 and 0.5 for $X_{O,0} = 0.21$. The dilution and temperature effects were isolated in additional experiments with manipulation of N₂/Ar ratio in the oxidizer side, similar to [568]. The data shown in Figure 31, with fixed temperatures, clearly demonstrate the notable effects of concentration modification on soot formation. Note, in Figure 31, the SVF was plotted against convective time from the flame front to account for the fuel stream inert addition that would alter the flame position, and thus the residence time of soot particles. A further analysis of the slope of SVF with time indicated that the soot production rate varied linearly with initial fuel concentration $X_{F,0}$, provided that the flame temperature remained constant.
Figure 31. Separation of temperature and dilution effects on soot formation in ethylene CDFs, data taken from Axelbaum et al. [74].

As previously discussed, sooting limits in CDFs can be represented by the critical strain rate \( K_{cr} \), beyond which soot scattering signals disappear at a fixed \( X_{F,0} \) and \( X_{O,0} \) [551]. Because of the effects of inert addition in inhibiting soot formation, it may also be represented by the critical \( X_{F,0} \) and \( X_{O,0} \), below which soot cannot form altogether at a fixed strain rate. Indeed, a CDF can be manipulated to be sooting or non-sooting by varying the amount of \( N_2 \) dilution in the fuel or oxidizer stream. As mentioned before, a sooting map can be drawn in the \( X_{F,0} - X_{O,0} \) plane, and a further example (obtained by Joo et al. for ethylene fuel [281]) can be seen in Figure 32a. The critical sooting conditions (black squares) in terms of \( X_{F,0} \) and \( X_{O,0} \), were determined from scattering signals where soot signals are about to appear, similar to the description in [551]. In subsequent studies, similar soot maps were obtained for various other fuels and fuel mixtures [614, 620], which were then used to compare the sooting tendency of these neat, or mixture fuels.
Figure 32. Sooting limit curve of counterflow diffusion flames as determined with light scattering technique (a), and with visual inspection (b). Data taken from Refs. [281] and [581] for (a) and (b), respectively.

Instead of using light scattering, Naik et al. [581] determined the critical sooting condition based on visual inspection of the flame, so that the first observation of yellow soot luminosity against a darkened background was deemed to be a signal for incipient soot formation. Figure 32b shows the soot map, developed with this visual criterion for methane CDFs at a constant global strain rate $K_G$ of 20 s$^{-1}$. A notable difference from light scattering-based measurements lies in the higher $X_O$ region, where the obvious bending behavior, observed in the light scattering-based sooting limit curve (a) is absent in the visually-based curve (b). Although the two curves were measured for different fuels, later light scattering-based experiments [620] confirmed that the phenomena of bending in sooting limit curves are ubiquitous, and the bending is even stronger for low sooting tendency fuels like methane.
Before explaining this discrepancy, it is helpful to identify several important features exhibited by the sooting limit curve shown in Figure 32a: 1) Beginning from the condition of $X_{F,0} = 1.0$, the critical $X_{O,0}$ for soot onset increases only slightly as $X_{F,0}$ decreases down to a value of about 0.4, beyond which the critical $X_{O,0}$ increases sharply. Such dramatic variations indicate that sooting behaviors in the large and small $X_{F,0}$ regions are rather different, caused by the transitions of flame/sooting structures. This is consistent with the distinction between SF and SFO flames, mentioned previously; 2) In SF flames with large $X_{F,0}$, the sooting limits closely follow the iso-temperature curve, suggesting that incipient soot formation is controlled by temperature. On the other hand, in SFO flames with low $X_{F,0}$ and large $X_{O,0}$, sooting limits span an appreciable range of iso-temperature curves, implying that temperature is not the only factor controlling soot formation in SFO flames; 3) The most interesting observation is, in fact, the bending behavior of the sooting limit curve of Figure 32a, which can be crossed twice by a vertical line of constant $X_{F,0}$ (e.g., the line of $X_{F,0} = 0.22$, not drawn). This has important implications in that for flames with a fixed $X_{F,0}$, as $X_{O,0}$ increases, the initial non-sooting flame can have a transition to the sooting regime and then become non-sooting again as $X_{O,0}$ approaches unity, exhibiting a non-monotonic behavior with $X_{O,0}$. Indeed, soot volume fraction measurements and accompanying numerical modelling results confirmed this non-monotonic influence of oxidizer composition on soot formation in CDFs with low $X_{F,0}$ [378]. In particular, it was experimentally shown that for ethylene CDF with $X_{F,0} = 0.28$ and an effective strain rate $K_E$ of $100 \text{s}^{-1}$, the peak SVF first increased—reaching a maximum near $X_{O,0} = 0.7$—and then decreased as $X_{O,0}$ increased from 0.7 to 1.0 [378]. Further kinetic analyses revealed that, while increasing $X_{O,0}$ suppresses soot inception and enhances soot oxidation (tending to reduce soot production), soot surface growth rates are enhanced with the increase in $X_{O,0}$, contributing positively to soot mass concentration. It is these competing effects that lead to the non-monotonic variation of the sooting tendency with $X_{O,0}$ in SFO flames. Since the visually-based sooting limit curve shows no sign of bending (implying simply a monotonic dependence), it can be concluded that visual inspection may
not be suitable for judging the onset of soot—at least for CDFs of SFO types. This is consistent with the findings of Du et al. [551], that the sooting limits determined by visual luminosity can be strongly affected by fuel type and flame geometry, and with Zhang et al. [525], who postulated that gaseous PAHs may also be responsible for the yellow color emissions in CDFs.

Compared to argon and nitrogen, helium has a special feature, that of high diffusivity. Since species diffusion is always coupled with chemical kinetics to determine general flame characteristics, it is reasonable to expect that the soot formation process would also be noticeably affected by preferential diffusion [701-703]. In this regard, the effects of helium addition on the sooting characteristics of diffusion flames have attracted wide research interests. For instance, McLintock [704] compared the effectiveness of adding various inert gases in an oxidizer stream in reducing the sooting propensity of ethylene coflow diffusion flames, and found helium to be most effective. However, in cases of fuel stream addition, Glassman and coworkers [705] noticed that helium was even less effective than argon in inhibiting soot formation of ethylene coflow flames. Axelbaum et al. [657] proposed an explanation for these interesting opposing trends based on concentration modification: When added to the fuel side, preferential diffusion effects will cause helium to defocus to a higher extent than argon, due to the concave curvature of the coflow diffusion flame front (i.e., negative stretch) [659]. As a result, concentrations of the fuel and other fuel-related intermediate species in the helium-diluted flame will be higher (effectively less diluted) than those in the argon-diluted flame, leading to higher soot formation in the helium-diluted flame. The opposite holds when the diluents are added to the oxidizer side, so that helium will be more focused, resulting in more effective dilution, and lowering the sooting tendency in the helium-diluted flame.

In addition to the effects of concentration modification, helium has a higher thermal diffusivity, so its addition can also cause a broadening of the temperature profile, which may thermally affect soot formation. By using a positively stretched flame, for which temperature broadening has the opposite effects (if any) to that of concentration modification on soot formation, Axelbaum et al. [657]
provided unambiguous evidence to support the dominance of concentration modification as the underlying mechanism explaining the helium addition effects. In particular, noble gases such as helium, argon, neon, and krypton that have similar heat capacities, but progressively lower diffusivities, were added to the fuel stream (at a molar ratio of 0.2) of positively stretched CDFs. The results (Figure 33) showed that peak SVF was lowest in the case of helium addition for both ethylene and propane flames. Since temperature broadening due to the high thermal diffusivity of helium would tend to increase soot formation if effective, the experimental facts can only be explained by composition modifications. Note that the relative extent of the effects between fuel stream helium and argon dilutions were expected to be reversed as the signs of stretch rates changed. This is supported by later experimental investigations [706], where the fuel stream addition of hydrogen (a highly mobile gas, similar to helium) was found be less effective than N₂ addition in reducing soot production in ethylene coflow flames, which can also be partly explained by this mechanism, especially considering that H₂ has an additional chemical role in inhibiting soot formation in ethylene flames [662].
Figure 33. Effects of inert dilution (20% by mole) on soot volume fraction profiles in CDFs of C₂H₄ (a) and C₃H₈ (b). Data taken from Axelbaum et al. [657].

5.2.2 Carbon dioxide and water vapor addition

Although typically regarded as non-reactive, CO₂ and H₂O can participate in chemical reactions under high temperature flame conditions, so they could chemically affect fuel oxidation and pollutant formation processes. Detailed investigations on the effects of CO₂ and H₂O addition on soot formation can shed light on the underlying mechanism of these intricate chemical interactions (and help validate and improve existing kinetic models). Besides, the frequent use of exhaust/flue gas recirculation (EGR/FGR) techniques in practical combustion devices [694, 707, 708] also motivates such studies.

Du et al. [564] experimentally determined the critical strain rate $K_{cr}$ for soot inception in ethylene and propane CDFs and found that soot formation was inhibited by CO₂ addition in either the fuel or oxidizer side, for both flames. They also showed that CO₂ addition inhibits soot formation via a combination of dilution, thermal, and chemical effects; although they did not identify detailed
reaction pathways that are chemically affected by the addition of CO$_2$. Zhang et al. [525] compared the experimentally measured C$_2$H$_2$ mole fraction, SVF, and soot number density between a baseline ethylene CDF and one with 21% CO$_2$ addition on the fuel side, which have the same $X_{F,0}$ and $X_{O,0}$ and similar flame temperatures (via reactant preheating for the CO$_2$ addition case). The data showed that CO$_2$ addition resulted in notable reduction of C$_2$H$_2$ mole fraction and SVF, attributed (through logical reasoning by the authors) to the fact that CO$_2$ addition enhanced the reaction of CO$_2$ + H = OH + CO to produce more OH radicals in the fuel pyrolysis zone, which then contributed to the oxidation of gas phase soot precursors. Annunziata et al. [709] measured the laser-induced emission (excited at 266 nm) of ethylene CDFs and found the emissions detected both at 330/440 nm (termed LIF signals by the authors to represent nanoparticles constituted mainly by aromatic compounds) and at 550 nm (termed LII signal to represent soot aggregates of sizes higher than 10 nm) were significantly reduced by the addition of CO$_2$, either in the fuel or the oxidizer side, which were primarily attributed to the higher heat capacity of CO$_2$, thereby the thermal effects.

To further clarify the chemically-inhibitive role of CO$_2$ in soot formation, Liu et al. [710] performed numerical simulations with a detailed chemistry for ethylene CDFs. An innovative strategy was employed in the simulation, which used fictitious CO$_2$ (FCO$_2$), defined purely for numerical purposes, as a species that has the same thermodynamic/transport properties with real CO$_2$ but does not participate in any chemical reactions. Chemical effects of CO$_2$ addition were then identified by comparing the results obtained with FCO$_2$ and those from with real CO$_2$. The results indicated that the reduction in flame temperature, C$_2$H$_2$ concentration, and enhanced oxidative attacks by O and OH in the sooting zone, via CO$_2$ + H = OH + CO and CO$_2$ + CH = CO + HCO, were the underlying mechanisms for the chemical role of CO$_2$ in reducing soot formation. It was also found that adding CO$_2$ to the oxidizer side was more effective in inhibiting sooting tendencies than in the fuel side. However, this study did not directly model PAH/soot formation, instead it used the changes of C$_2$H$_2$ mole fraction as an index to represent sooting tendency variations. Using the same FCO$_2$ strategy,
Wang and Chung [626] performed PAH-based soot modelling studies to determine the effects of fuel side CO₂ addition on ethylene CDFs with detailed gas-phase chemistry and the method of moments to treat particle dynamics. After validation against the experimental data, numerical results showed that the addition of CO₂ had chemical effects in: 1) reducing PAH mole fraction, which in turn decreased soot nucleation rates and number density, and thus lower surface area for mass-adding reactions, 2) decreasing hydrogen, methyl, and propargyl radicals, causing lower H abstraction rates in the production of active surface sites for soot growth, and 3) reducing C₂H₂ concentrations, and thus the HACA growth rates per unit surface area. They further measured the sooting limits in terms of critical \( X_{F,0} \) and \( X_{O,0} \) in CO₂ atmosphere (i.e., both the fuel and oxygen are balanced by CO₂) and the results are shown in Figure 34, with comparison against data measured in N₂ atmosphere. It can be seen that the sooting region in the \( X_{F,0} - X_{O,0} \) plane was appreciably reduced in the case of CO₂ dilution. In addition, when plotted against computed flame temperature, it is clear that for SF flames (e.g., \( 1000/T_f [K] > 0.5 \)) at a fixed \( X_{F,0} \), the flame temperatures at the sooting limit condition were higher in the CO₂ atmosphere than those in N₂. Furthermore, the temperature differences between the CO₂ and N₂ diluted flames were also higher for lower \( X_{F,0} \) conditions (i.e., more dilution), providing further evidence of the chemical effects (in addition to temperature effects) of CO₂ addition on soot inception.
Figure 34. Sooting limit curves of C$_2$H$_4$ counterflow diffusion flames (a) and $X_{F,cr}$ and $X_{O,cr}$ as a function of calculated flame temperature (b) for C$_2$H$_4$ CDFs with N$_2$ and CO$_2$ dilutions. Data taken from Wang and Chung [626].

The above investigations in non-premixed flames consistently showed that CO$_2$ addition can reduce soot formation, however, it is of interests to mention that there are also studies that indicated the opposite. For instance, Abian et al. [707, 711] observed that a 25% CO$_2$ addition in ethylene actually increased soot yield during experiments on temperature-controlled flow reactor fuel pyrolysis, although higher levels of CO$_2$ addition led to reduced soot formation. The authors explained the enhancement of soot formation with a low level of CO$_2$ addition, based on the increased OH radical concentrations through CO$_2$ + H = CO + OH, which were expected to increase consumption of H$_2$ by OH + H$_2$ = H$_2$O + H. The removal of H$_2$ will favor H abstraction reactions in the HACA mechanism for PAH/soot growth and thus leads to higher soot production. At a higher dilution level, CO$_2$ may result in a relatively large amount of OH radicals, which help to oxidize PAHs and soot particles. Similar enhancement of soot by CO$_2$ was seen by Teini et al. [712] in
experiments of C₂H₂ pyrolysis. Further kinetic simulation showed that the addition of CO₂ enhanced the production of C₃H₃ (and thus benzene by C₃H₃ self-combination) by increasing OH radicals, which were also seen to help abstract H atoms from PAH molecules, allowing for further HACA growth. Note that in these pyrolysis experiments, the thermal effects of CO₂ were not present because temperatures were held fixed in all cases, and the reactants are O₂ deficient. In this way, these observations of CO₂ increasing soot formation share many similarities with the finding that a small amount of O₂ addition in the fuel side of C₂H₄ [165] and C₂H₂ [713] fuels actually enhanced soot formation. This is especially true considering the fact that CO₂ addition was not found to enhance soot production during the pyrolysis of methane [712], consistent with Gülder’s findings in methane coflow flames [714]. This will be discussed in more detail in a later section on partial premixing.

Compared to CO₂, there are much fewer studies on the effects of water vapor addition on soot formation in CDFs, probably due to the additional experimental complications of metering and vaporizing liquid water. Equipped with a counterflow burner with the capability of reactant preheating, Zhang et al. [525] found that the addition of water vapor to both the oxidizer and fuel side could reduce soot formation, although addition on the oxidizer side was more effective. It was also qualitatively noticed that water vapor addition was more effective for soot inhibition than adding the same moles of CO₂. Although no detailed kinetic simulation was performed, the authors reasoned that, with H₂O + H = OH + H₂, the added water produced considerable amounts of OH radicals, which helped oxidize molecular soot precursors. The higher effectiveness of H₂O compared to CO₂ was explained simply by the fact that more OH radicals can be produced with H₂O addition, as a consequence of the faster rate of H₂O + H = OH + H₂ than CO₂ + H = OH + CO [715]. On the other hand, experimental results, as obtained with coflow diffusion flames, led Schug et al. [705] to conclude that H₂O plays only a thermal effect. A similar conclusion was drawn by Rao and Bardon [716], who burned liquid hydrocarbon/water emulsions in diffusion flames. In an effort to clarify whether H₂O can chemically affect soot formation, Liu et al. [717] conducted numerical simulations
of sooting coflow diffusion flames of ethylene with oxidizer side H₂O addition. Computations with fictitious H₂O addition were included and it was observed that the chemical effects of H₂O led to higher flame temperatures, but lower soot concentrations. Specifically, the addition of water reduced O radical concentration via O + H₂O = OH + OH, so that the concentration of CH₂, produced primarily from C₂H₂ + O = CH₂ + CO, was also reduced. This explained the chemical effects of H₂O in reducing the formation of aromatic soot precursors such as benzene and pyrene, because CH₂ is essential in the formation of C₃H₃ (via e.g., C₂H₂ + CH₂ = C₃H₃ + H), the self-recombination of which is the major benzene formation pathway. Furthermore, through H + H₂O = OH + H₂, the concentration of H radical was also reduced, decreasing HACA growth rates.

5.2.3 Partial-premixing

Partial-premixing refers to the addition of oxygen—either pure or diluted by inert gases—in the fuel side of a non-premixed combustion system. The level of addition is typically kept low so that the equivalence ratio of the premixture is much larger than unity in the fuel stream. Motivation for studying the effect of partial premixing on soot formation includes the fact that a small amount of oxygen is frequently present in the fuel side of diffusion flames. As an example, for coflow flames which are widely adopted in studying soot formation, a reaction quenching occurs in regions in close proximity to the nozzle due to excessive heat and radical losses, so that a small amount of air entrainment is inevitable [109]. Oxygen penetration into the fuel side can also occur due to finite rate chemistry (i.e., incomplete combustion) [660]. From a practical point of view, air is sometimes intentionally co-injected with fuel in industrial combustion devices, to improve fuel atomization. It was also shown that in diesel engines ahead of the main diffusion flame zone a fuel-rich premixed region is always present influencing initial soot formation typically due to a lifted nature of flame base [718, 719]. Furthermore, studies on partial-premixing may also clarify the underlying
mechanisms that can help identify dominant soot/PAH growth pathways in flames of different fuels. Notable examples can be seen, for example, in the work of McEnally and Pfefferle [720].

A previous section presented the general sooting structures of partially-premixed CDFs; a more detailed and quantitative discussion on the hydrodynamic and chemical effects of partial-premixing on soot formation is given here. It is helpful to first show some direct images of partially premixed CDFs. Figure 35 presents the variation of the visual flames as an incipiently-sooting non-premixed counterflow flame (equivalence ratio $\phi = \infty$) transited towards partially-premixed flames up to $\phi = 3$ (taken by Carbone et al. [81]). Note here that the stoichiometric mixture fraction ($Z_{st}$) and peak temperature were held constant as progressively more oxygen was added to the fuel stream, in order to maintain the location of the diffusion flame sheet and the thermal structure unchanged on the oxidizer side. It can be seen that—apart from the purely non-premixed flame—all the other partially-premixed flames (PPF) featured double flame zones (represented by the CH chemiluminescence, in blue) with the bottom zone being rich premixed flame and the upper zone diffusion flame. The soot zones (yellow luminosity) were sandwiched between these two flames. With the aid of thermocouples for temperature measurement, and GC-MS for species quantification, Carbone et al. [81] performed careful experimental investigations to resolve the detailed chemical structures of these flames. It was shown that oxygen addition led to a much more rapid temperature rise as the fuel stream was convected downstream, as a result, there was more time for soot-related reactions to occur in the high temperature regions. In fact, increases in C$_3$, C$_4$ (except for C$_4$H$_2$), benzene, and soot concentrations were found as $\phi$ decreased. Further kinetic analysis also revealed that an enhanced contribution from C$_3$ pathways to benzene formation was induced by oxygen addition. It should be mentioned that the lowest level of premixing tested by Carbone et al. [81] was already high enough ($\phi = 6.5$) to generate a separate rich-premixed flame. As seen later, there are other studies in which the level of premixing was much lower and no separate second flame can be observed. In these cases, the mechanism of
partial premixing on soot formation can be different. In subsequent discussions, these cases will be termed as *small* amounts of oxygen addition.

![Figure 35](image_url)

*Figure 35.* Direct images of partially-premixed ethylene counterflow flames with decreasing equivalence ratio in the fuel side, reprinted from Carbone *et al.* [81] with permission of Elsevier. Fuel and oxidizer streams issued from bottom and top, respectively. Yellow line indicates position of stagnation plane, and the flame with $\phi=3$ had a lower temperature than the rest.

Hura and Glassman [113] are among the pioneers to measure soot volume fraction in CDFs with partial premixing. A comprehensive investigation was performed to test the effects of small amounts of fuel side oxygen addition on soot formation in CDFs of ethylene, propene, propane, n-butane, and iso-butane [113]. The results showed that soot volume fraction, average particle sizes, and surface growth rates were all noticeably increased with an addition of 10% oxygen to the ethylene flame (a doubling of peak SVF); while the enhancing effects were rather small for propene flame, with even suppressive effect observed in propane and iso-butane flames. The authors noticed that oxygen addition in ethylene led to an increased amount of H radical, which helped to promote fuel pyrolysis rate. However, no details were reported regarding important soot precursors. Realizing that fuel side oxygen addition may increase flame temperature, and thus thermally affect soot formation, Hura and Glassman, in a subsequent study [721], manipulated the oxidizer composition of the CDF to keep the peak flame temperature unchanged. They also extended the range of oxygen addition from an equivalence ratio of infinity (purely diffusion flame) to 3.0 and 2.6 for ethylene and propane flames, respectively. Major results of this study are re-plotted in *Figure 36* in terms of the variation of peak extinction coefficient (i.e., an indication of SVF) with equivalence ratio. Similar soot enhancing/inhibiting effects were observed for small amounts of oxygen addition (large equivalence
ratio, to avoid significant variations in flame structures) in ethylene (propane) flames. With further addition of oxygen, both flames exhibited a sharp increase, followed by a rapid decrease in soot formation, attributable to the drastic change of flame structures. While enhanced fuel pyrolysis rates explained the initial soot increase in ethylene, fuel dilution was believed to be the dominant factor in decreasing soot production in propane CDF.

Figure 36. Peak extinction coefficients for ethylene and propane partially premixed counterflow diffusion flames as a function of fuel stream equivalence ratio, $f_w$ is fuel flow rate. Reprinted from Hura and Glassman [721] with permission of Elsevier.

The fact that small amounts of oxygen chemically enhance ethylene sooting tendency was confirmed by a number of other investigations in CDFs [165, 564] and coflow diffusion flames [564, 705, 714, 722-727]. The influence of oxygen in propane flames, on the contrary, has been the subject of much debate, and disagreement appeared at two levels. First, contradictory results were reported in different studies regarding the overall effect of fuel side oxygen addition on soot formation. For instance, Du et al. [564] showed that the critical strain rate $K_{cr}$ (i.e., sooting tendency) of propane CDF increased monotonically with the level of oxygen addition, up to 50%, while Hura and Glassman [113] observed a decrease in peak SVF when 10% oxygen was added. Similarly, in coflow diffusion flames of propane, increased soot productions were seen in [564, 725, 728], but reduced soot formation was observed in [721, 722, 724] with a small amount of oxygen addition. No definite
explanation was found to explain these discrepancies. The second level of disagreement concerned the chemical effects of oxygen addition. Even though the phenomenon of soot reduction observed by Hura and Glassman [721] was qualitatively consistent with the experimental results of Gülder [714], explanations for the reduction differed. Specifically, through reactant preheating to maintain adiabatic flame temperature and comparison against the N₂-diluted flames, Gülder experimentally separated the chemical effects of oxygen addition from thermal and dilution effects. His results showed that oxygen significantly enhanced soot formation through its chemical interactions [714]. However, Hura and Glassman [721] concluded that oxygen was mainly a fuel diluent, with insignificant chemical effects, and Du et al. [564] showed a significantly suppressive role for oxygen in propane flame. Note that although the work of Hwang and Chung [165] showed a reduction of SVF in propane CDFs with oxygen addition, in their study the oxidizer that comprised both oxygen and nitrogen was added to the fuel stream (to maintain temperature), so that the observed soot reduction was a combined effect of chemical and dilution, and it was unclear whether or not soot formation would be suppressed if allowance was made for the dilution effects. It appears that additional experiments—especially those accompanied by detailed kinetic simulation—are required to clarify this matter.

Based on a detailed gas-phase kinetic simulation [165], the enhancement of soot formation by oxygen addition in ethylene CDFs was explained through the increased reaction rates of C₂H₂ + O = CH₂ + CO. Methylene (CH₂) is essential for the formation of C₃H₃—thus the first aromatic ring. With on-line mass spectrometry measurements, McEnally and Pfefferle [68] experimentally observed (and later confirmed by numerical simulations [729, 730]) that peak concentrations of CH₄, C₃H₃, and 1,3-C₄H₆ increased with the addition of oxygen (in the form of air) in the fuel stream of ethylene coflow diffusion flames, providing support for the C₃H₃-based explanation in [165]. (Note that 1,3-C₄H₆ can be formed with the addition of CH₃ to C₃H₃.) In addition to the above-mentioned reaction for methylene production, it was also shown that oxygen addition can enhance C₂H₄ + O = CH₃ + HCO to produce CH₃, which is then involved in various C₁ + C₂ interactions to enhance C₃ species
formation. As mentioned in previous discussions regarding the synergistic effects of C_2H_4-based binary fuel mixtures, C_2H_4 flames are particularly sensitive to increased C_1 species from either fuel dopants or oxidative pyrolysis, because it is somewhat difficult to break the strong C=C double bond at pyrolysis temperatures [713], which leads to a deficiency of the C_1 species. Following this reasoning, similar phenomenon can be expected in C_2H_4 diffusion flames, considering the strength of the C≡C triple bond. In fact, in the C_2H_2 CDFs studied by Leusden and Peters [75], small amounts of added oxygen were seen to chemically enhance the formation of C_3H_4, benzene, and soot. Further kinetic analysis showed that both the C_3 and C_4 + C_2 channels for benzene formation were promoted. Similar results were also noticed in coflow diffusion flames of C_2H_2 [713]. Since 1,3-butadiene (CH_2=CH-CH=CH_2) shares a feature with C_2H_4/C_2H_2 in that its pyrolysis does not easily produce C_1 species, it may be expected that oxygen addition would also increase soot formation in 1,3-butadiene flames. However, experimental data in coflow diffusion flames showed the opposite to be true. It was postulated by McEnally and Pfefferle [713] that 1,3-butadiene may isomerize to CH_2=C=CH-CH_3 so that C_1 and C_3 species can be easily produced by C-C bond breaking. Alternatively, the dominant benzene formation channels for 1,3-butadiene may be through even carbon chemistry (making C_3 chemistry less important), which has been partially confirmed by a recent chemical speciation study [174].

Partial-premixing is also interesting for methane flames, since natural gas appliances typically rely on partially-premixed flames. Mungekar and Atreya [561, 562] studied the effects of partial-premixing on soot formation in low strain rate methane CDFs, for both the SF and SFO configurations. For SF flames, the authors observed that with increasingly higher levels of oxygen addition, the soot zone thickness first decreased (from φ = ∞ to 8.3) and then increased (from φ = 8.3 to 6.4). While the initial decrease (consistent with the observations of Nakamura et al. [731] in a methane CDF with φ decreasing from infinity to 10) was thought to be caused by the shrinkage of the distance between the non-premixed flame front and the stagnation plane, it was presumed that the subsequent increase was
due to the presence of a separate high-temperature rich premixed flame front, on the fuel side of the non-premixed flame front, where soot began to form. Peak SVFs were observed to decrease as $\phi$ decreased from infinity to 8.3; this was explained based on the dilution and chemical suppressive role of O$_2$, which was also shown to contribute to the soot reduction in methane coflow diffusion flames with partial-premixing [714, 732]. As $\phi$ further decreased, the high temperature non-premixed flame front moved closer to the stagnation plane, leading to increased temperature and longer soot residence time in high temperature regions, and resulting in an increase in SVF. In SFO flames, where the flame is located on the fuel side of the stagnation plane, fuel stream partial premixing was seen to monotonically decrease soot formation; while it was noted that such a decrease was consistent with the competition between soot inception, growth and oxidation, no detailed explanations were provided.

Although without soot measurements, Berta et al. performed temperature and intermediate species measurements for pre-vaporized n-heptane non-premixed flames [592] and PPFs [563, 584] in a counterflow configuration, with special focus on gaseous soot precursors, including C$_2$H$_2$ and benzene. The oxidizer streams in their flames were air in all cases, while the fuel stream consisted of n-heptane vapor diluted with various amounts of air, producing various equivalence ratios. The structures of the PPFs were resolved both experimentally and numerically, with satisfactory agreements. The axial profiles of temperature, velocity, C$_2$H$_2$, C$_2$H$_4$ and C$_6$H$_6$ are shown in Figure 37 for various equivalence ratios. Results showed that as $\phi$ decreased, the diffusion flame front tended to move closer to the fuel nozzle, consistent with the work of Mungekar and Atreya [561, 562], and much as expected, considering the variations in stoichiometric requirements. The rich premixed flame zone (indicated by the bump in the velocity profile on the fuel side of the diffusion flame) was also seen to move to the fuel nozzle as $\phi$ decreased, which can be rationalized by considering the balance between the stretched laminar flame speed and the incoming reactant velocity. As $\phi$ decreased from infinity, the flame speed of the pre-mixture increased so that the flame front moved to a location at a
higher reactant speed, i.e., closer to the fuel nozzle. The stagnation plane, on the contrary, moved away from the fuel nozzle as $\phi$ decreased. An increased level of partial-premixing was seen to monotonically reduce C$_2$H$_2$, C$_2$H$_4$ C$_6$H$_6$, as well as coronene (C$_{24}$H$_{12}$), and the effects were much stronger for large PAHs (mole fractions for coronene, C$_6$H$_6$ and C$_2$H$_2$ decreased by a factor of around 1,000, 10 and 2, respectively, as $\phi$ decreased from 15 to 2.5 [563]). This was attributed by the authors to reduced residence time and enhanced oxidation. This result contrasts with the observations of Mungekar and Atreya [561] in methane PPFs, where an increase of soot was observed as $\phi$ decreased from 8.3 to 6.4. The major difference was that while Mungekar and Atreya [561] partially-premixed the fuel with pure oxygen, Berta et al. [563] added air in the fuel stream so that a large amount of N$_2$ was present in the fuel stream with partial-premixing. In these cases, significant fuel dilution may have been the dominant contributor to the observed soot reduction.

![Figure 37](image)

**Figure 37.** Axial profiles of temperature and axial velocity (a), and C$_2$H$_2$, C$_2$H$_4$ and C$_6$H$_6$ mole fractions (b) of partially-premixed counterflow diffusion flames at various equivalence ratios. Data taken from Berta et al. [563].
5.2.4 Other dopants

a) addition of chlorinated species

Interest in halogenated species originated with their use as flame extinguishing agents [733-735], primarily due to their ability to scavenge the highly reactive H radicals in hydrocarbon flames [736]. There were also concerns about environmental pollution and the problems of incineration-based disposal of hazardous wastes [737], of which chlorinated species (as in chemical solvents, polychlorinated biphenyl (PCB) and polyvinyl chloride) may account for a considerable fraction.

Many investigations in shock tubes [738] and premixed flames [739, 740], as well as diffusion flames [741], have shown that chlorinated compounds can enhance PAH/soot formation, as compared to their hydrocarbon counterparts. For instance, a systematic work of Senkan et al. [739] showed that the critical equivalence ratio ($\phi_c$) for soot formation decreased in a methane-based premixed flame with the addition of CH$_2$Cl$_2$, CHCl$_3$, and CCl$_4$, and the effects were stronger for molecules containing more chlorine atoms. A critical review of relevant studies until the early 1990s was provided by Morse and Cundy [742]. Subsequent works included the development of detailed kinetic models to understand fundamental combustion characteristics of chlorinated hydrocarbons (CHC) [420, 743-745]. Several possible reasons were provided to explain the higher sooting tendencies of CHCs: First, as the C-Cl bond is noticeably weaker than C-H bond [746], initial fuel pyrolysis is easier for CHCs, which can lead to higher concentrations of the radicals essential for PAH/soot growth [737, 738, 740]; Second, the chlorine atom enhances H-abstraction reaction from stable PAH molecules through $A_r + Cl \rightleftharpoons A_{r'} + HCl$, leading to more aromatic radicals for further HACA growth [737]; Third, chlorine may also scavenge oxidative radicals like O and OH, resulting in reduced soot oxidation [740], although such effects are not present in oxygen-free pyrolysis experiments.

Relevant studies conducted in CDFs are somewhat scarce. Yang and Kennedy [569] performed an experimental investigation of CDFs fueled with pure CH$_4$ and CH$_4$/CH$_3$Cl binary mixtures, with air
as the oxidizer, and at a constant local strain rate $K_L$ of 38.7 s$^{-1}$. The neat CH$_4$ flames did not produce measurable amounts of soot, as a result, only the soot volume fraction profiles of the more sooting CH$_3$Cl-doped flames were measured. The data showed that the sooting structure of CH$_3$Cl-doped flame was qualitatively similar to a typical hydrocarbon SF CDF. However, quantitatively, peak SVF increased monotonically–and almost linearly–with the molar ratio of CH$_3$Cl in the CH$_4$/CH$_3$Cl mixtures. Leylegian [594] measured the critical sooting strain rate ($K_{cr}$) in CDFs of CH$_4$/CH$_3$Cl and C$_2$H$_4$/CH$_3$Cl binary mixtures and found a nearly linear increase of $K_{cr}$ with a molar mixing ratio of CH$_3$Cl, a result that was qualitatively consistent with that of Yang and Kennedy [569]. An interesting phenomenon was observed for C$_2$H$_4$/CH$_3$Cl mixture in that $K_{cr}$ first increased and then decreased, exhibiting a non-monotonic variation trend with CH$_3$Cl mixing ratio. The author believed this was caused by chemical interactions between the mixture components, i.e., the initial increase can be attributed to the effect of chlorine on the fuel pyrolysis rates, while the latter decrease was due to the lowered ethylene concentration. However, it is important to point out that in order for the author to maintain the carbon density (in g/cm$^3$) constant in the fuel stream, the N$_2$ dilution (mole fraction) was higher in cases with lower CH$_3$Cl mixing ratio, so that the different dilution levels may also play a role.

b) addition of metal-containing compounds

Investigations on combustion of hydrocarbon fuels doped with metal-containing compounds are generally related to the following topics: 1) flame inhibition [747-749]; 2) synthesis of metal oxide nanoparticles [750-752] and desired carbon nanostructures (i.e., carbon nanotubes) [753-755]; 3) PM/soot emission reduction [118, 756-759]. Here we are concerned only with soot, and it was found that iron-containing compounds such as ferrocene and iron-pentacarbonyl are studied most.

Several mechanisms have been proposed to explain the effects of metal additives on soot formation: First, some metals, such as sodium and potassium, can easily become ionized in flame environments.
Their charges can subsequently be transferred to soot precursors so that the rates of soot nucleation are effectively reduced [118]. Although more evidence is needed, this mechanism is known to be consistent with observations of rich premixed ethylene flames where potassium chloride (KCl) doping decreased both the soot volume fraction and the particle size [756, 760]; Second, metal containing compounds can scavenge the active radicals of H, O and OH, affecting the fuel pyrolysis processes [748], through which PAH/soot formation can, in turn, be influenced; Third, metal-containing compounds—especially iron-containing species—can decompose to form metal nanoparticles which serve as the nuclei to provide additional surface sites for soot growth [761]; Fourth, metal-containing species, when incorporated in soot particles, can notably catalyze soot oxidation [263]. Note that the third and fourth mechanisms contribute in opposite ways to the final soot emission, while both are concretely supported by data in the literature. For instance, shock tube studies showed that soot yields for C₃H₈/Ar and C₂H₂/Ar mixtures can be significantly enhanced after the addition of a small amount of iron pentacarbonyl (Fe(CO)₅) [762]. Studies in rich premixed flames, where a deficiency of oxidizing species limits soot oxidation, have shown enhanced soot formation after doping of either ferrocene or Fe(CO)₅ [761, 763, 764]. It was also observed that in diffusion flames, particles appeared earlier in iron-seeded flames than in unseeded flames [262, 263, 634, 765]. These studies supported the third mechanism, as discussed above [757]. Strong evidence also exists for the fourth mechanism, both for in-flame measurements near the oxidizing flame front [263, 757], and ex-situ oxidation measurements of iron-containing soot [766]. In particular, the oxidation activation energy for soot formed in iron-containing environments was found to be 116 kJ/mol, significantly lower than the 162 kJ/mol for soot formed in unseeded flames [767]. Furthermore, a recent study suggested that the iron-containing particles that catalyze soot oxidation were primarily elemental Fe, rather than any form of iron oxide [768]. It is generally accepted that soot reduction by metal addition (observed in many practical combustion devices) is a result of more efficient burnout, rather than inhibition of soot nucleation [761].
Relevant studies in CDFs were performed by Raj et al. [261, 634, 769] who doped Fe(CO)$_5$ in a methane CDF with $X_{Fe}=0.097$, $X_{O}=0.53$ and an effective strain rate $K_E$ of 17 s$^{-1}$. Of special note is that Fe (CO)$_5$ was carried by a separate N$_2$ stream and fed to the flame through a central tube embedded in—and concentric with—the bottom fuel burner nozzle. As a result, the flame was not quasi-1D, as in conventional CDFs. The authors observed a significant decrease in the concentrations of C$_2$ species (i.e., C$_2$H$_6$, C$_2$H$_4$ and C$_2$H$_2$) with Fe (CO)$_5$ addition and attributed these reductions to the catalytic combustion of methane by iron oxide [261]. This explanation was consistent with the observed reduction of CO and faster consumption of CH$_4$, while it seemed incompatible with the data of Kim and Hahn [768], which indicated the dominance of elemental iron over iron oxide of the diffusion flames. Nevertheless, in a later modelling work of these flames, Raj et al. [769] mentioned that the reduction of C$_2$ species could be due to scavenging of O, H and OH radicals by iron species; while these radicals could have contributed to molecular growth from methane to C$_2$ species. Perhaps the most striking observation of Raj et al. was the presence of large amounts of O$_2$ in the fuel side for the seeded methane CDF, even at locations very close to the fuel outlet [261]. The authors explained this interesting behavior by the diffusion of oxygen across the stagnation plane, due to early consumption of the fuel; unfortunately, however, no oxygen concentration profiles were provided in a subsequent modelling study [769] for further confirmation. Raj et al. [634] continued to study the particle evolution of the same CDF with and without Fe (CO)$_5$ addition. With tube sampling of the flame gas and subsequent particle analysis with a SMPS, the total particle number density and the size distributions were measured at different axial positions between the two opposing nozzles. Major results are shown in Figure 38, where it can be seen that under catalytic conditions, particles appeared earlier near the fuel side, and the peak soot number density was higher than in the unseeded case. However, the peak average particle diameter was smaller for the seeded flame, suggesting that the addition of Fe (CO)$_5$ resulted in the formation of larger numbers of smaller particles. The reduced C$_2$H$_2$ concentration [261] (thus slower HACA growth), and possibly reduced coagulation rates in the
seeded flame, may help rationalize such findings. More interesting is that both the soot number density and average size in the seeded flame were seen to decrease more rapidly towards the flame front, offering evidence for the catalytic effect of iron-containing species on soot oxidation, and consistent with previous studies [261].

\[\text{Figure 38. Particle number concentrations (a) and average particle diameters (b) of CDFs with and without fuel-side iron pentacarbonyl addition. Note the axial profiles along both the axis of symmetry (R = 0 mm) and along a radially shifted vertical line (R = 5 mm) are shown, considering the influence of the central tube for precursors. Adapted from Raj et al. [634] with permission of Elsevier.} \]

It is noted that there are other dopants, such as sulfur- and nitrogen-containing compounds (i.e., H\textsubscript{2}S, SO\textsubscript{2}, SO\textsubscript{3}, NH\textsubscript{3}, NO and NO\textsubscript{2}) which have been tested for their effects on soot formation [119]; however, since no relevant studies were performed in CDFs, these dopants will not be discussed further here.
5.3 Effects of pressure on soot formation

Energy conversion systems such as IC engines, jet engines and gas turbines typically rely on combustion at high pressure for maximum thermal efficiency and energy-releasing intensity, making it of practical interests to study soot formation at high-pressure conditions. Understanding the dependence of soot formation kinetics on pressure is also of vital importance from a scientific point of view. Experimental investigations of soot formation in high-pressure flames are notably more difficult than those performed at atmospheric conditions, partly because of flame stability issues [107] and partially due to the additional complexities of apparatus and instrumentation [127]. Also, the reduced flame thickness and excessive soot formation at high-pressure conditions can cause significant difficulties for quantitative soot measurements. Nevertheless, many researchers have successfully established steady flat premixed flames [770] and coflow diffusion flames [20] at pressures up to 100 atm.

A review of early studies on pressure effects was provided by McArragher and Tan [771]. Subsequently, optical methods such as color pyrometry and laser light extinction / scattering were frequently applied in investigations of sooting characteristics in high-pressure flames. For premixed flames, focus was on the variations of final SVF, soot growth rate, particle number density and concentrations of important gaseous intermediates with pressure [770, 772-777]. In particular, at a fixed equivalence ratio and temperature, the final SVF were seen to increase with pressure ($P$) proportional to $P^2$ up to several bar (approximately 7 bar) [772], beyond which a linear relationship between SVF and pressure was observed, indicating the absence of further soot mass growth at those extremely high-pressure conditions [770]. Later numerical calculations confirmed the observed pressure dependence [778] and the decay of soot growth was found to be caused either by decreasing particle activity at moderate pressure (15 bar) [774], or the lack of a growth agent (i.e., $C_2H_2$), at even higher pressures (70 bar) [770]. Similarly, the effect of pressure on the radical levels, especially those of H radicals, which plays an important role in PAH and soot growth, can be notable. It was also
observed that pressure increase tended to reduce particle coagulation rates, attributed to the alteration of soot surface properties in high-pressure conditions that affect particle-particle sticking [776].

Soot formation in high-pressure diffusion flames has received more attention in the last decade, justifiable by the dominance of non-premixed combustion in soot-emitting practical combustion systems. A comprehensive review of soot studies in laminar diffusion flames before 2012 was provided by Karatas and Gülder [107], in which coflow flames has been investigated to the greatest extent. The advantage of coflow diffusion flames with concentric circular nozzles is that the flame heights can be maintained reasonably independent of pressure, provided that the fuel mass flow rate (i.e., in mg/s) is fixed [107, 779, 780]. Also, at a given height above the fuel outlet, the average velocity does not vary much with pressure because the increase in flow density is balanced by the decrease in the flow cross-section area [483, 781]. As a result, soot volume fraction can be directly compared at the same height to flames under different pressure conditions, providing tractability of the pressure effects. From the review by Karatas and Gülder [107], it is noted that before 2012, flames of gaseous fuels accounted for the majority of these studies. At a constant fuel mass flow rate, increases in pressure generally led to a narrowing of the flame, resulting in increasingly curved inwards. Peak SVF, line-integrated SVF, and maximum conversion of fuel carbon to soot, all increased with pressure in the form [soot] \( \propto P^n \), where \( n \) was fuel dependent [61] and became smaller in the higher end of the pressure ranges [483]. The increase of soot yield resulted not only from flame narrowing, but from overall soot formation enhancement. Computational studies [782, 783] suggested that the pressure increase results in earlier soot inception, but later oxidation, thus experiencing more growth time. It was also shown [784] that high pressure forces coflowing air to infiltrate into the fuel side near the flame base, resulting in enhanced fuel pyrolysis.

More recently, research interest has gradually shifted towards pre-vaporized liquid fuels such as ethanol [785], C6 hydrocarbon and oxygenated species [21], n-heptane [486, 786, 787] and n-decane [445]. The effects of pressure on soot properties other than SVF were also tackled, using light
scattering [69] and/or thermophoretic sampling [509]. This section is intended to complement the review on coflow high-pressure flames in 2012 [107] by examining soot formation in high-pressure CDFs. In fact, Figura and Gomez [788] showed that CDFs are largely immune to buoyancy-driven instabilities, while this may pose significant issues for high-pressure coflow diffusion flames, especially those with elongated flame shapes, and thus, higher Richardson numbers. Together with its advantages of quasi-1D structures, clean boundaries and easy control of residence time, CDF is believed to be an ideal flame reactor for soot studies in high-pressure non-premixed systems. Nevertheless, upper constraints in the pressure range for steady CDF are caused by the transition towards turbulent flames, although the use of helium as a diluent may delay such transition [788].

The sooting limits represented by critical strain rate $K_{cr}$ were experimentally determined by Du \textit{et al.} [573] for ethylene-air CDFs in the pressure range from 1 to 2.5 atm; the results are summarized in Figure 39. Similar to previous studies at atmospheric pressure [551], laser light scattering was used to identify critical sooting conditions; and partial substitution of N$_2$ with Ar in the oxidizer stream helped maintain the peak flame temperature as the pressure was varied. As can be seen, $K_{cr}$ increased monotonically, indicating an enhanced sooting tendency as pressure increased (a). Also as expected, at a specified pressure, $K_{cr}$ decreased with the decrease of $X_{F,0}$, demonstrating the effects of dilution on soot formation. The authors argued that the effects of pressure would be better represented by density-weighted critical strain rate, considering that, instead of the purely kinematic strain rate $K$, the characteristic reaction rates in CDF are proportional to the density-weighted strain rate $\rho_0 K$ [789] (where $\rho_0$ is the density of the oxidizer at the nozzle exit). Interestingly, a linear relationship was observed between $\rho_0 K_{cr}$ and pressure $P$ (b), and it was further shown that the data collected at different $X_{F,0}$ collapsed into a single line in the form of $\rho_0 K_{cr} \sim X_{F,0}^{1/2} P$ (c). Accompanying numerical simulation showed that the maximum partial pressure of C$_2$H$_2$ was also linearly correlated with $X_{F,0}^{1/2} P$, so that $\rho_0 K_{cr} \sim P_{C2H2}$. With these results, the authors concluded that the soot formation rate in the investigated C$_2$H$_4$ flames was of first order with C$_2$H$_2$ concentration [573]. In a follow-up study [85], CH$_4$ and
C\textsubscript{3}H\textsubscript{8}–air CDFs were tested with an extended pressure ranging from 1 to 5 atm. The correlation between $\rho_0K_{cr}$ and $X_{F,0}^{1/2}P$ was again found to hold. When plotted against $P_{C2H2}$, as shown in Figure 40, the $\rho_0K_{cr}$ data for C\textsubscript{3}H\textsubscript{8} and C\textsubscript{2}H\textsubscript{4} flames collapsed into a single straight line, while the data for the CH\textsubscript{4} flame followed a distinctively different line. The authors rationalized the similarity between C\textsubscript{3}H\textsubscript{8} and C\textsubscript{2}H\textsubscript{4} flames by pointing out that C\textsubscript{2}H\textsubscript{4} was among the primary products of fuel-side propane pyrolysis. Sarnacki and Chelliah [530] performed absolute irradiance-calibrated time-resolved LII and PIV measurements to quantify SVF and flow fields, respectively, in ethylene CDFs up to 30 atm. While direct comparisons of soot loadings in flames of different pressures were difficult because different dilution/strain rates were used as pressure varied, the authors showed that global activation energy for nucleation increased exponentially with pressure up to 20 atm, beyond which a sudden decrease was observed. A correlation between the density-weighted critical sooting strain rate and $X_{F,0}^{1/2}P$ was confirmed again (with the exception of the data point beyond 20 atm), although a methodology different from previous studies was used to determine the critical strain rate [85, 573].
Figure 39. Effects of pressure on the critical strain rate (a) and density-weighted critical strain rate (b) for soot formation in counterflow diffusion flames. Also shown is the linear correlation between $\rho_0 K_{cr}$ and $X_{F,0}^{1/2} P$ (c). Reprinted from Du et al. [573] with permission of Elsevier.
The effects of pressure on PAH formation were investigated by Böhm and Lacas [575] for methane-air CDFs from 1 to 6 bar with $X_{F,0} = 1.0$ and with a global strain rate $K_G = 200 \text{ s}^{-1}$. PAH concentrations were estimated experimentally by LIF signal intensities (excited at 283 nm and detected at around 315 nm) and determined computationally by simulations with detailed chemistry. An approximate square dependence was observed for the peak PAH LIF signal with the increase of pressure (LIF intensity $\sim p^2$), consistent with the calculated maximum PAH concentration (summing up to pyrene) and previous experimental data obtained in premixed $C_2H_4$ flames. It should be pointed out that the flames studied by Böhm and Lacas [575] were close to the sooting limit but essentially non-sooting. Note that the increase in pressure also led to an increase in computed peak flame temperature (2000 K at 1 bar, 2100 K at 3 bar and 2150 K at 6 bar). Thus, the authors commented that the higher PAH concentration at elevated pressures was not only an effect of pressure. It is
relevant that the increase in peak temperature with pressure was also experimentally confirmed by Figura and Gomez [788] in methane CDFs using the technique of SiC thin filament pyrometry [790]; and it was shown that the increase can be more than the change of adiabatic flame temperature with pressure [636].

Comprehensive chemical speciation in high pressure CDFs was pioneered by Figura and Gomez [618]. In particular, with micro-probe sampling and subsequent GC-MS analysis the authors measured the axial profiles of major species, C2-C5 intermediate hydrocarbons as well as aromatic species, up to indene. The choice of flame conditions was based on the consideration of flame stability, adiabaticity, pressure tractability, and sooting level. Too much soot was undesirable as it would cause micro-probe clogging during the sampling process, while non-sooting flame would have rather low concentrations of PAHs, which was of special interest to the authors. Two series of flames at incipient sooting conditions were finally chosen for experiments, as well as numerical simulations using a detailed reaction mechanism accounting for PAH formation [791]. Flame series one featured a constant fuel mass fraction ($Y_{F,0}$) of 0.122, a stoichiometric mixture fraction ($Z_{st}$) of 0.408 and a global strain rate of $K_G = 57 \text{ s}^{-1}$, while the pressure was varied from 1 to 8 bar. Flame series two covered only two flames at 8.55 and 25 bar, with $Y_{F,0} = 0.0975$, $Z_{st} = 0.408$ and $K_G = 18.4 \text{ s}^{-1}$. The reduction of $Y_F$ in the second series was intended to avoid excessive soot loading at the high-pressure conditions (25 bar). A reduction of micro-probe outer diameter ($d_{out}$) was shown to be necessary for probing CDF at increasingly higher pressures, if sufficient spatial resolution was to be achieved [623]. In fact, it was suggested that $\delta/d_{out}$ should be kept above 20 [623], where $\delta$ is the thickness of the mixing layer, which decreases as pressure increases. In flame series one, with increased pressure, the experimental peak mole fractions of CH4, C2H6, C3H8, C3H6, n-C4H10 and 1-C4H8 decreased and those of 1,3-C4H6, C4H4, C4H2, c-C5H6, methyl-cyclopentadiene, and all tested aromatic species, increased, while the C2H2 mole fraction remained comparable. The enhancements in sooting tendencies from increasing pressure in both flame series were evidenced by significant increases in indene mole
fractions at the highest pressure conditions; this was attributed by the authors to the increases in reactant concentrations with pressure [618]. Besides providing quantified concentration data to assist kinetic model developments for high-pressure applications, an additional contribution of the work was an extensive discussion on scaling and the invariance of temperature-time history under different pressure conditions. The authors demonstrated that for a fixed strain rate, the mixing layer thickness $\delta$ of the CDF was proportional to $P^{-1/2}$. After normalizing the axial position and temperature by $\delta$ and peak temperature, respectively, the non-dimensionalized temperature profiles for flames at different pressures (but the same $Y_{F,0}$, $Y_{O,0}$ and $K_G$) collapsed into a single line. It was further shown that the temperature-time history within the thermal layer would be invariant for flames at different pressures, provided that these flames have a constant strain rate. Such invariance provides tractability in comparing flames operated at different pressures, otherwise, variations in residence time would add to the pressure effects to an unknown extent [618, 623].

In a more recent study [636], the density-weighted global strain rates ($\rho K_G$), instead of strain rates $K_G$, were kept constant for moderately sooting ethylene CDFs ranging 1-8 atm. SVF profiles were measured with LII excited at 532 nm, and detected at around 450 nm, calibrated against measurements with light extinction. Nozzle exit velocities (and thus $K_G$) had to be progressively lower to counteract the pressure increase to maintain a fixed $\rho K_G$. As expected, both peak local and integrated SVF increased with increased pressure. The pressure scaling factor $n$ was derived from the experimental data by plotting $\ln \left( f_{v,max}/f_{v,max,0} \right)$ as a function of $\ln \left( P/P_0 \right)$ and taking the slope, where the subscript 0 refers to a reference pressure state (e.g., at 1 atm). It was found that the value of $n$ varied with $X_{F,0}$ and $X_{O,0}$, and for a fixed pair of $X_{F,0}/X_{O,0}$, $n$ was larger in cases with constant $\rho K_G$ than in cases with constant $K_G$. The authors also showed that the thickness of the soot zone and the thermal mixing layer became invariant with pressure, as long as $\rho K_G$ was fixed, consistent with previous scaling arguments [618]. Based on this fact, the authors pointed out the importance of maintaining $\rho K_G$—rather than $K_G$—at a constant value to study pressure effects. However, it may be worthwhile to point out that
with $\rho K_G$ fixed, the variation of flow residence time with pressure may complicate the isolation of pressure effects, as demonstrated by Figura and Gomez [618, 623].

Amin and Roberts [529] pioneered in-situ determination of the morphology of soot particles in CDFs at elevated pressures. Specifically, laser light extinction and two-angle scattering were used to measure SVF, primary particle size, and number density, along with aggregate size (average radius of gyration) and number density in ethylene CDF at 2-5 atm with $X_{F,0} = 0.3$ and $X_{O,0} = 0.21$. Note that $K_G = 30 \text{s}^{-1}$, rather than $\rho K_G$ was kept constant. A special high-pressure vessel with four curved windows was built, providing $160^\circ$ of optical access to allow collection of scattering signals at various angles with respect to the incident beam. The results are summarized in Figure 41. Skewed profiles of SVF $f_v$ (a) and scattering coefficient $K_{VV}$ (b) can be seen, consistent with soot evolution in SF flames. With the increase in pressure, SVF (a), primary particle size $d_p$ (c), and aggregate size $R_g$ (d) were all increased, but the primary particle number density (c) decreased. The reduction of soot number density was explained by enhanced coagulation efficiencies, and it is noted that a pressure increase was reported to reduce coagulation efficiency in premixed flames [776]. Peak soot yield was found to be scaled with pressure raised to the power of 2.67 [529].
Carbone et al. [628] measured the mole fraction profiles of major and critical soot related intermediate species in C\textsubscript{2}H\textsubscript{4} counterflow partially-premixed flames at 1, 4, and 8 atm. Three fuel side equivalence ratios (∞, 6.5 and 5.0) were tested at each pressure, while keeping Z\textsubscript{st} and K\textsubscript{G} constant for all the flames. For consistent temperature-time histories for the reactants (so that the effects of pressure could be better isolated), the peak flame temperatures at high-pressure flames were deliberately reduced by increasing dilution. Nevertheless, the benzene mole fraction was still seen to increase for partially-premixed flames with increasing pressure, despite the higher dilution level. Reduced back diffusion of OH radical was thought to be a contributor to the enhanced soot precursor formation, aside from the pressure effects in increasing reactant concentrations [618]. Through kinetic
simulations, it was observed that with increasing pressure, the \( C_2 + C_4 \) pathways for benzene formation became increasingly more important than the \( C_3 + C_3 \) pathways.

More recently, Gleason et al. [490] complemented the experimental database on chemical structures of high-pressure counterflow flames with soot volume fraction measurements. Flame conditions were carefully selected to make sure that the normalized temperature-time history was identical, regardless of pressure. The measurement result showed that soot volume fraction increased by two orders of magnitude as pressure increased from 1 to 4 atm while peak flame temperature was kept unchanged. Further analysis on soot growth rate also demonstrated that the activation energies for high-temperature soot growth are the same at different pressures.

This discussion on pressure effects concludes by affirming the great necessity for more quantified PAH and soot characterization in high-pressure CDFs, with well-controlled conditions, especially for practical liquid fuels and their surrogates. Both challenges and opportunities exist for investigations on how, and to what extent pressure-dependent reactions affect soot formation, a topic that has been largely unexplored by existing studies.

5.4 Effects of temperature on soot formation

The qualitative and quantitative dependence of soot formation on temperature has received extensive research attention, justifiable by the fact that soot formation is a highly temperature-sensitive process. Temperature is a major influence not only on the formation of soot particles, but also on their oxidation. Early relevant investigations generally focused on the effects of temperature on critical C/O ratios \((C/O)_{cr}\), or critical equivalence ratios \((\phi_{cr})\) for soot appearance in premixed flames [98, 685, 772, 778, 792-795], and sooting height, or SVF profiles, in jet diffusion flames [96, 728, 796, 797].

For burner-stabilized premixed flames, it was generally observed that \( \phi_{cr} \) first decreases (i.e., increase of sooting tendency) and then increases with the progressive increase in temperature,
exhibiting an interesting non-monotonic variation [772]. At a constant C/O ratio, SVF was also seen to peak at a certain temperature, and either an increase or a decrease from this temperature would decrease soot formation [97, 772, 794]. These findings can be partially understood by the competing effects of temperature on soot precursor formation and on oxidative destruction. In the low end of the temperature range, an increase in temperature would result in faster fuel decomposition to soot precursors and an enhancement of the molecular growth processes towards PAHs and soot particles. However, further increase in temperature would result in a faster increase in the oxidation rates of soot precursors/particles than in the rates for their formation [796]. Also, reduced surface reactivity [597] and decreased thermodynamic stability of soot precursors [141] can play important roles in reducing soot formation at high temperatures.

For jet diffusion flames, Glassman and coworkers [96, 796] showed that sooting height decreases with the increase in adiabatic flame temperature, demonstrating a qualitative trend in these flames that increases in temperature lead to an enhancement of the sooting tendency. Note that the adjustment of flame temperature was achieved by the authors through inert dilution (N₂ or Ar) in the fuel stream, which may affect soot formation through both flame temperature and concentration modifications. Nevertheless, the authors proved that the effect of temperature is dominant and fuel dilution plays only a secondary role. Although this conclusion seems to be consistent with the sooting limit curve obtained for SF CDFs (e.g., Fig. 26, with relatively large X₅) where temperature dominates the critical condition of soot formation, in other cases it was reported that dilution can have important and even greater effect than temperature [95, 700].

In diffusion flames, soot inception and growth occur on the fuel side of the flame, where the concentration of oxidizing species is rather low. Without the competing effect of oxidation, an increase in flame temperature is expected to enhance the rates of soot precursor formation, as well as those of particle inception and surface growth, leading to a monotonic increase of sooting tendency. Note that for over-ventilated coflow diffusion flames, oxidation occurs later in the post-flame zone.
as heterogenous reactions, which will notably affect ultimate soot emission, but not the critical sooting condition [796]. Also note that fuel pyrolysis in diffusion flames is not the same as in a shock tube, where it was shown that soot yields have non-monotonic variation with temperature [798]. In a shock tube, pyrolysis occurs at uniform temperature, and if this temperature is too high, the thermodynamic stability of soot precursors decreases, and soot production is reduced [141, 798]. On the other hand, there is always a spatial profile of temperature in diffusion flames, extending from boundary temperatures in the fuel/oxidizer inlets all the way to peak flame temperature. Fuel pyrolysis and soot inception can always be efficient locally at a location with intermediate temperatures. As a result, the sooting tendency in diffusion flames generally increases monotonically with temperature. Beltrame et al. [576] showed that in a soot formation CDF with undiluted methane as the fuel stream, peak SVF increased monotonically as $X_{O,0}$ increased from 21% all the way to 100% (although it should be noted that increases of $X_{O,0}$ reduce dilution as well as increasing temperature).

In fact, the variation of temperature of diffusion flames was frequently achieved by adjusting the amount and type of diluents in both fuel and oxidizer streams, so that the effects of dilution were likely to interfere with the effects of temperature. Therefore, dilution effects must be isolated if a quantitative correlation between soot formation and temperature is to be obtained. In this regard, Axelbaum et al. [74] adjusted the flame temperature of CDFs by the substitution of volume flow rate of $N_2$ in the oxidizer stream and an equal volume flow rate of Ar. Since Ar has a lower heat capacity than $N_2$, its substitution resulted in an increase in peak flame temperature. Since the mole fraction of the inert diluent was not affected, temperature could be varied without affecting dilution level. Using such an approach, the effects of dilution and temperature can be separated. In practice, flames with different levels of fuel stream dilution can be compared at a fixed peak temperature; and flames with different peak temperatures can also be compared at constant $X_{O,0}$ and $X_{F,0}$. Results from these types of studies were discussed in a previous section on inert dilution and will not be repeated here.
A well-controlled peak temperature does not necessarily mean a controlled spatial temperature profile [74], and the latter may be of importance for the sooting process in SF CDFs, where the particle growth region is spatially separated from the peak flame temperature region. Ideally, the effects of temperature on soot formation should be studied with a well-controlled, well-characterized temperature-time history (i.e., not only the peak temperature) which the soot particles may experience. Towards this goal, Carbone et al. [628] demonstrated the importance of keeping the stoichiometric mixture fraction ($Z_{st}$) and the global strain rate ($K_G$) constant when perturbing peak flame temperatures. By doing this, the position of the flame front relative to the gas stagnation plane—as well as the flame thickness—can be kept constant so that after normalizing the temperature with respect to its peak value, the normalized temperature-convective time history remains similar among flames with different peak temperatures. In other words, the temperature-time profiles become self-similar, so the effects of peak temperature could be effectively isolated [489, 490]. Note, in order to achieve peak temperature variation while keeping $Z_{st}$ constant, the dilution levels in both the fuel and oxidizer streams must be altered, although this additional complexity caused by the variations of dilution level may be insignificant if the required temperature range is moderate. In particular, Gomez and coworkers [489] showed that in order for the peak temperature of ethylene CDF to vary from 1890 to 2186 K (variation of around 300 K), the fuel mass fraction must increase from 0.30 to 0.39, while the oxygen mass fraction in the oxidizer stream must increase from 0.23 to 0.30. The measured axial profiles of temperature, SVF and soot production rate for this series of ethylene CDFs are shown in Figure 42.
As can be seen in Figure 42 (a) and (b), although the temperature profiles are self-similar, the shape of the SVF profiles vary among the different peak temperature cases. In these flames, the SVF begins to increase in a region close to the flame front and then continues to grow as they are convected towards the particle stagnation plane, exhibiting common features for SF flames. However, the rates of soot growth vary notably among the flames. As shown clearly in Figure 42 (c), for cases with lower peak temperature, the initial growth rates are also lower. Since this initial growth occurs in a high

Figure 42. Axial profiles of temperature (a), SVF (b), and soot production rates (c) for ethylene CDFs with well-controlled convective time-temperature history. Adapted from Carbone et al. [489] with permission of Elsevier.
temperature region \((Z \sim 0.5 \text{ mm})\), where HACA surface reactions are expected to be efficient, their decreasing rates, with decreasing peak temperatures, are consistent with the high activation energy characteristics of the HACA reaction scheme. As particles are transported towards the stagnation plane, excessive temperature reduction causes the soot growth rates to fall quickly to nearly vanishing values. A somewhat unexpected observation was a sharp increase in soot growth rates in regions very close to the stagnation plane, where the temperature was below 1600 K. The authors suggested a possible ion-induced nucleation mechanism to explain this. Although this seems to be qualitatively consistent with the modelling results of Wang et al. [140], where nucleation reaction and physical addition of PAH were seen to contribute to soot growth in the proximity of the particle stagnation plane, the comparable peak level of this low-temperature growth rate with that of the high-temperature HACA rate was not predicted by Wang et al. More studies, involving both numerical and experimental efforts, are needed to shed light on the mechanisms of this interesting low-temperature soot growth. It is also important to point out that Sirignano et al. [89] also found evidence of low temperature soot growth and demonstrated that soot particles can also exist in the fuel side of the particle stagnation plane.

5.5 Effects of strain rate on soot formation

In Section 4.3, we discussed how strain rate may influence the overall sooting structure of CDFs. Here we intend to complement that discussion with a more comprehensive review on strain rate effects on PAH and soot formation. Because PAH / soot formation is a kinetically-driven rate-limiting process, quantitative investigations on the correlation between sooting characteristics and strain rate in CDFs can help explain the kinetics of soot formation.

A second motivation for studying strain rate effects is the link between diffusion flamelet and turbulent flame, the latter is somewhat difficult to model due to the large variations of temporal and spatial scales involved. The task becomes even more challenging when soot formation chemistry is
considered, which by itself includes a large number of reactions, spanning a wide range of reaction time scales. The flamelet model, as pioneered by Peters [106], provides a tractable approach for simulating turbulent diffusion flames with complex chemistry. In such a model, local structures of the turbulent reaction zones are approximated by strained flamelets, which are to be convected and stretched by the turbulent flow fields. A key parameter characterizing diffusion flamelets is scalar dissipation rate ($\chi$), closely related to, and with the same unit (reciprocal of time) of strain rate. In fact, the structures of steady flamelets can be solved entirely in the mixture fraction space once $\chi$ is specified [799]. In this context, a CDF is an appropriate model because of its relevance to many physics postulated for diffusion flamelets. The strain rate of CDF can be linked to $\chi$ of diffusion flamelet in a straightforward way. Besides, efforts needed to systematically vary the strain rate of CDF are minimal as it can be achieved by simply adjusting the nozzle exit velocities. This is in contrast to coflow flames, in which the strain rate is difficult to control and usually too small to be a representative of turbulent flames [114].

It is important to highlight here that due to the slow kinetics involved in PAH/soot formation, sooting process in highly transient flames can have significant unsteady effects. The above-mentioned (steady) flamelet approximation is not able to predict the formation of soot, although it may be used with success in predicting the overall local structure of turbulent flames (i.e., major flame products). As a result, it is of practical interests to investigate soot formation under unsteady strain rate conditions. Section 5.5.2 is dedicated to this matter.

5.5.1 Strain rate effects on sooting behaviors in steady CDFs

A number of studies have investigated how strain rate affects PAH formation, both experimentally [91, 114, 585] and computationally [91, 92, 114, 800]. The experimental techniques that were used to evaluate PAH concentrations were either gas chromatography or LIF; while chromatography can provide quantified mole fractions of individual PAH species, LIF can only provide fluorescence
intensities that qualitatively represent a class of PAHs in a certain size range. These studies generally demonstrated that as the strain rate increases, peak flame temperature, C₂H₂ and C₃H₃ decrease moderately, and the mole fractions of aromatic species decrease more significantly. In addition, PAH species with larger molecular sizes are more sensitive to strain rate. An example is shown in Figure 43 for C₂H₄ CDF with X_F,₀ = 1.0 and X_O,₀ = 0.25 [91], where both numerical and experimental data are provided. A similar trend was also observed in a study of CH₄–air CDF, where the investigated strain rate spanned a much larger range (i.e., K_G from 100s⁻¹ all the way to 600 s⁻¹, close to extinction) [114]. More quantitative GC speciation analysis [590] of a C₂H₂ CDF with X_F,₀ = 0.375 and X_O,₀ = 0.220 (balanced by Ar) showed that, as the effective strain rate K_E increased from 38 to 50 s⁻¹, the peak mole fraction of benzene (A1) decreased from 2×10⁻⁴ to 8×10⁻⁵, by approximately 40%. This is to be compared against naphthalene (A2, 4×10⁻⁵ to 5×10⁻⁶) and pyrene (A4, 3×10⁻⁵ to 4×10⁻⁷), the mole fractions of which exhibited a decrease of one and two orders of magnitude, respectively.

Figure 43. Effects of strain rate on PAH formation in ethylene counterflow diffusion flames. Both computed peak PAH mole fractions and experimental peak PAH LIF intensities are shown. Note PAH LIF signals detected at larger wavelengths are expected to represent PAHs with larger molecular sizes. Reprinted from Wang and Chung [91] with permission of Elsevier.

The effects of strain rate on flame and PAH chemistries result from several mechanisms. First, the flame intensity (and thus peak flame temperature) are expected to decrease with the increased strain rate, primarily because of the reduced Damköhler number. Since the temperature-dependent fuel
pyrolysis process is essential in diffusion flames to produce small molecular precursors (i.e., C2H2, C3H3, C4H3, etc.) for PAH growth, decreased temperature would lead to lower PAH formation rate; Second, even with abundant small molecular precursors, the growth of PAH is a slow rate-limiting process and would be significantly inhibited by decreased residence time. Since larger PAHs grow sequentially from smaller ones, this also explains why larger PAHs are progressively more sensitive to strain rate; Third, the general scaling argument [81] shows that the increase in strain rate would reduce the characteristic thickness of the flame, resulting in the originally-separated pyrolysis zone and high temperature OH-oxidizing zone being pushed closer together. As a result, concentrations of oxidizing species in the fuel pyrolysis zone tend to increase, causing increased fuel oxidation to form sequentially CH2O, CHO and CO. Note, however, this last mechanism is expected to be important only at very high strain rates (i.e., close to extinction) [114].

The first quantitative measurement of the variation of SVF with strain rate was conducted in CDFs stabilized around a cylindrical porous burner using laser light extinction and scattering [554]. Similar experimental studies were later performed in counterflow burners with opposing screened, or contoured, nozzles [78, 91, 576]. Much as expected, these studies generally showed that the peak SVF exhibited a monotonic decrease as the strain rate increased, as has already been discussed in Section 4.3.

It is logical to expect that fuels with distinct molecular structures will have different sooting rates when burned under similar conditions. For instance, it is widely accepted that benzene formation is a critical step, for being rate-limiting, toward the formation of soot in flames of small aliphatic fuels. On the other hand, the situation is obviously different for fuels that already contain aromatic rings (benzene formation can be bypassed). Similar arguments can be applied even to different fuels belonging to the same class, considering that different fuel molecules lead to pyrolysis/oxidative intermediate species that contribute to soot precursor formation through different channels and at different rates. Variations in the sooting rate of different flames are expected to lead to variations in
the sensitivity of the final soot concentrations to strain rate. Decroix and Roberts [801] found the rates of decrease in peak SVF to be higher for C$_3$H$_8$ than in C$_2$H$_4$ CDF. Similar experimental results were obtained by Wang and Chung [91], with additional measurements for C$_3$H$_6$ and C$_2$H$_6$ flame. Their results showed that the peak SVF in C$_2$H$_6$ flame had a higher sensitivity than C$_3$H$_8$ flame to strain, and C$_3$H$_6$ flame had the lowest sensitivity. In addition, numerical simulations [92] showed that the dependence of peak SVF to strain rate can be fitted into a power law relation: $f_{v,max}(K_G)/f_{v,max}(K_{G0}) = (K_{G0}/K_G)^a$ where $K_{G0}$ is a reference global strain rate and the subscript $a$ is the exponent, with higher values indicating higher sensitivity of SVF to strain rate. The value of $a$ was found to be fuel-dependent, and it was 0.88 for benzene flame, much smaller than the value of 1.2 and 2.9 for ethylene and methane flames, respectively [92].

A more interesting observation [91] is that, although the sensitivity of soot loadings to strain rate depend on fuel type at a fixed flame condition ($X_F$,0, $X_O$,0), once the peak SVF is adjusted (for instance, by varying $X_O$,0) to be similar among these different fuels at a given reference strain rate, the sensitivity of peak SVF to strain rate becomes reasonably independent of fuel type. This point is clarified in Figure 44, where the correlation between peak SVFs and their slope of decrease with strain rate (represented by nozzle exit velocity $V_0$ in the figure) are shown. Although the data were collected for various fuels, they fall on the same line, demonstrating that the peak SVF at a reference strain rate (not fuel type) dominates the strain rate sensitivity. This interesting result seems to indicate an inherent similarity in the soot formation processes among different fuels. It should be remembered, however, that this conclusion is only drawn from the four aliphatic fuels tested in Figure 44, and further investigations are necessary to test whether it can be generalized to other fuels. It is also noted that this result is consistent with the sooting sensitivity index (SSI) data [620], developed to quantify the dependence of sooting tendency of a particular fuel to strain rate variations.
Figure 44. Correlation between soot loading and its sensitivity to strain rate. Reprinted from Wang and Chung [91] with permission of Elsevier.

In addition to global soot loading, as characterized by SVF, the effects of strain rate on the details of particle size distribution were also investigated. Numerical simulation with a sectional method [92] showed that in both benzene and ethylene SF CDFs, the percentage contribution of the number of small particles (~1.7 nm) to the total soot number density was only marginally affected, but those of larger particles decreased notably with the increase in strain rates.

5.5.2 Soot formation in CDFs subject to unsteady strain rates

The preceding discussions focused on sooting behaviors at steady conditions. For application in turbulent combustion modelling through the flamelet approach, unsteady effects are also important [802]. In a turbulent environment, the mean strain rate of flamelets is established from the large-scale eddies, while fluctuations around the mean values and associated unsteady effects can be caused by smaller eddies, with turnover time comparable to the diffusion time scale of the flamelets [803-805]. CDF is an ideal flame configuration to study unsteady strain rate effects because the unsteadiness can be established experimentally by imposing oscillation on the nozzle exit velocity. Numerically, the quasi-one dimensionality makes time-resolved simulation with detailed chemistry much more efficient.
Before addressing soot formation in unsteady CDFs, a general understanding of global flame responses to oscillating strain rates is helpful. Relevant studies on the dynamics of unsteady CDFs [804, 806-812] showed that for small oscillation frequencies of nozzle exit velocity (i.e., strain rate), the flame responds to the oscillations in a quasi-steady manner (characterized by heat release rates, temperature, or OH concentration profiles). As the oscillation frequency increases, some transient behavior (e.g., phase lag, reduced response) begin to develop. With further increase in oscillation frequency, the flame may fail to respond altogether. The cutoff frequency that differentiates between quasi-steady and transient responses was also observed to increase in cases with stronger burning [808]. An example of the variation of maximum flame temperature with oscillation frequency can be seen in Figure 45. Through a time-resolved simulation with detailed chemistry, Egolfopoulos and Campbell [805] provided a physical interpretation of the above observations: On either side of the reaction zone in a CDF, there is a hydrodynamic zone, followed by a diffusion zone in which diffusion is the dominant mechanism for transporting fuel/oxidizer toward—and heat away from—the reaction zone. Although the effects of oscillation in the nozzle exit velocity can be propagated nearly unaltered through the hydrodynamic zone, the oscillation can only be sensed indirectly by the flame through the effects that far-field disturbances induce on the diffusion processes. It was further shown that the exit velocity oscillation in CDF creates an oscillation of mass fraction of the reactants. In this regard, mass diffusion of the reactants towards the reaction zone is the dominant mechanism controlling the flame response to oscillating boundary conditions. The authors proposed an analogy between the unsteady CDF and the Stokes second problem, in which the fluctuating velocities induced by an oscillating plate in an initially stagnant flow (through momentum diffusion) become attenuated with the increase in the distance away from the plate (z) and the oscillation frequency (ω). The amplitude of velocity fluctuation in Stokes second problem is controlled by the non-dimensional Stokes parameter \( \eta_z = z(\omega/2\nu) \), where \( \nu \) is the fluid kinematic viscosity. Similarly, it was shown that the amplitude of peak temperature fluctuation in unsteady CDFs is controlled by \( \eta_K = \delta(\omega/2D)^{1/2} \approx \)
\[(\omega/2K_G)^{1/2}\], where the latter approximation is supported by the dependence of the diffusion zone thickness \(\delta\) on the diffusivity \(D\), and the global strain rate as \(\delta \approx (D/K_G)^{1/2}\). The flame response would decrease as \(\eta_K\) increased, either by increasing frequency or reduced strain, as shown in Figure 46, where the flame response of many different flames collapsed into a single line with \(\eta_K\). Physically, a decrease in the global strain rate can lead to increases in diffusion thickness so that the attenuation in flame responses are expected to be stronger. The increase in frequency would be accompanied by disturbances with shorter wavelength, causing larger gradients and diffusive fluxes—again dampening flame responses.

![Figure 45](image_url)

**Figure 45.** Oscillations in peak flame temperature of CDFs subject to oscillating strain rates at various frequencies (a) and frequency response of the amplitudes of maximum flame temperature and phase shift (b). Data taken from Egolfopoulos and Campbell [805].
With overall flame behaviors under unsteady conditions clearly explained by Egolfopoulos and Campbell [805], more recent studies focused on how PAH and soot formation are affected by unsteadiness [395, 577, 585, 593, 598, 630, 801]. The response of PAH/soot formation to strain rate oscillations does not necessarily follow that of the peak flame temperatures or heat release since: 1) PAHs/soot are typically formed in the fuel rich pyrolysis zone that lie inside the diffusion zone of CFDs, compared to the thin reaction zone where most heat release occurs; 2) The chemical time scale for PAH/soot formation is notably larger than the main heat release reactions; 3) large PAHs and soot particles have very small diffusivity compared to gas-phase species. Time-resolved in-situ SVF measurements in unsteady CDFs were first performed by Decroix and Roberts [801] using LII. The oscillation in strain rate was experimentally achieved by driving two loudspeaker diaphragms, installed respectively at the end-faces of two plenums; through each plenum, the fuel and oxidizer were introduced to the nozzle outlets. Sine-wave driving voltages were used so that the global strain rate ($K_G$) varied in a sinusoidal manner around a prescribed mean value. Experiments were done at four initial $K_G$ ranging from 15 to 90 s$^{-1}$, with unsteady forcing at frequencies from 25 to 200 Hz, and at both low and high amplitudes. Results showed that maximum soot SVF (global maximum in terms
of both temporal and spatial variation) for the unsteady flames was generally higher than their steady counterparts, this enhancement was seen to be most prominent in cases of low forcing frequency and high amplitude. The SVF responses were much attenuated with high-frequency oscillations. For example, an increase by a factor of 3.4 in peak SVF was observed for propane CDF oscillated at 25 Hz around a mean $K_G$ of 60 s$^{-1}$ while soot production became nearly insensitive for oscillation frequencies larger than 100 Hz. Furthermore, significant phase lag between the peak SVF peaks and strain rate troughs were noticed.

Because PAHs are the precursors of soot, their behavior in unsteady flames is particularly important for a better understanding of soot formation. For this reason, planar LIF was used to detect spatially- and temporally-resolved PAH concentrations for unsteady CDFs [585]. The LIF signals were detected at different wavelengths to represent PAHs of different size classes, with lower wavelengths corresponding to smaller PAHs. As expected, the results showed that PAH concentrations only responded to low frequency oscillation, becoming insensitive to oscillations beyond a critical value, which depends on PAH sizes and several other flame conditions like fuel type and mean strain rate. For larger PAHs, this critical frequency was lower, which was reasonable since the formation of larger PAHs requires the presence of smaller ones. However, an unexpected finding was that the critical frequency was smaller for flames with a higher mean strain rate. For example, with an initial strain rate ($K_L$) increasing from 23 to 44, and then to 74 s$^{-1}$, the critical frequency for large PAH responses decreased from 200 to 50, and then to 30 Hz. This is in sharp contrast to the scaling laws proposed by Egolfopoulos and Campbell [805] in which a normalized parameter $\eta_K = (\omega/2K_G)^{1/2}$ controlled the amplitudes of the flame responses (i.e., flame temperature) to the oscillating strain rate. According to the definition of $\eta_K$, the critical frequency would be larger for flames with higher $K_G$. A physical interpretation (mentioned above) is that the thickness of the diffusion zone would be reduced so that free-stream oscillation could propagate into the reaction zone with less damping in flames with a higher strain rate. The notable deviation between the response of the PAH
concentrations and this scaling law for overall flame properties (i.e., peak temperature) clearly demonstrates that diffusion-limited responses alone are not sufficient to explain the PAH behavior observed under unsteady conditions. The relatively larger time scale for PAH formation compared to that of the main heat release reaction is likely to play a role. It is noteworthy that although the authors in the original publication [585] attempted to correlate PAH with $\eta K$, the scatter of the data was fairly clear.

Besides the critical frequency, the amplitudes, symmetries and phase lags of the variations in PAH/soot concentrations subject to unsteady strain rates are also important. Figure 47 shows the peak, trough and mean values, as well as phase lags for the PAH mole fractions as a function of oscillating frequencies [593]. Clearly, the phase lags are always close to zero at low frequencies, indicating quasi-steady responses. However, the peak and trough values for PAH/soot concentrations are not symmetric around the mean values. The asymmetry becomes more significant for larger PAHs—and most notably for soot. More interestingly, the mean PAH/soot concentrations (averaged over an entire oscillating period) are higher than the steady state value, consistent with the experimental data of Decroix and Roberts [585]. The asymmetry and the resulting increase in the average PAH/soot concentrations in the unsteady flames may be attributed to the fact that PAH/soot formations are more sensitive to the variations in strain rate (and thus residence time) at lower strain rate instants. This is more or less consistent with the findings in steady flames. For example, Beltrame et al. [576] showed that the peak SVF of a CH$_4$-O$_2$ CDF decreased by around 50% as the strain rate increased from 20 to 40 s$^{-1}$, while a further increase of strain rate by 20 s$^{-1}$ only resulted in a decrease of peak SVF by 28%.
Figure 47. Detailed frequency responses of induced oscillation amplitude and phase-lag of soot and its precursors (benzene, pyrene and coronene). Reprinted from Cuoci et al.[395] with permission of Elsevier.

As oscillating frequency continues to increase, notable phase lag develops between PAH/soot concentrations and strain rates. The phase lag was also seen to be larger for species with a larger characteristic reaction time scale [395, 630]. Cuoci et al. [395] explained this by referring to the diffusion zone through which free-stream oscillations must pass to affect the reaction zone, and the species-dependent responses were attributed to the different diffusion time of each species. Following analysis by Egolfopoulos and Campbell [805], the authors also tried to correlate the predicted oscillation amplitudes of PAH/soot concentrations with the relevant Stokes number; however, it was not possible to collapse the responses of different species into a single line. This was expected since only diffusion-limited responses were taken into account in the relevant discussions based on and in the very definition of the Stokes number, while the effects of slow PAH/soot chemistry were excluded. In fact, this point was considered by Rodrigues et al. [630] and it was shown analytically (through
linearized asymptotic analysis) that the phase lag and the reduction of response amplitudes of a certain flame quantity (i.e., species concentrations) with the increase in frequency, are directly affected by the Damköhler number of the corresponding flame quantity. In addition, the authors [630] also demonstrated that the steady response of any flame property to strain rate also affects its responses (i.e., amplitude damping) under unsteady conditions. It can be generalized that flame responses to strain rate oscillation are primarily determined by: 1) the Stokes number, which compares the characteristic diffusion time scale with the imposed oscillation time scale; 2) the Damköhler number, which indicates the characteristic reaction time scale of a certain quantify of interests; and 3) a parameter that represents the steady response of a certain quantity of interests to strain rate variations [630].

5.6 Various other effects

Noteworthy counterflow-based soot studies for various other effects are discussed next.

5.6.1 Soot formation in counterflow flames of fuel sprays

In addition to use in the study of flames with gas-phase or pre-vaporized fuels, counterflow burners have also been employed in spray combustion studies [813, 814]. Spray flame is a highly complex multi-phase phenomenon and much of its fundamental physics are still not fully understood. As such, experimental [815-817] and numerical [818-821] investigations of spray flames established in a simple flow field are useful for clarifying their major features. Previous studies have shown that droplet mean size, size distributions, and droplet evaporation rates are important factors that affect the general characteristics of spray flames, such as flame temperature and species concentration distributions [822-825]. Depending on the boundary conditions, both premixed-like and diffusion-like flames can exist in the combustion of fuel droplets [826]. Drop group combustion (i.e., the combustion of droplet clusters) was observed to behave like diffusion flames [827-829].
With respect to soot formation in spray counterflow flames, Chen and Gomez [816] studied two CDFs of quasi-monodispersed n-heptane sprays with the same global fuel/air ratios. The first flame existed as a thin blue sheet and was observed to behave similarly to that of a purely gaseous flame, which was rationalized by the fact that the droplets are completely evaporated before reaching the flame zone. In the second flame, however, with a higher initial mean droplet size and velocity, some droplets survived and penetrated the first flame of fuel vapors towards the oxygen-rich environment. As a result, a second flame, characterized by a thick orange region on the oxidizer side, was formed. The authors attributed this orange luminosity to radiation of small soot particles, although quantification of soot concentrations was not performed. Nevertheless, this study indicated that an increase in initial droplet size may enhance soot formation, a topic for which Hayashi et al. [603] conducted detailed experimental and numerical investigations. The authors established n-decane (n-C10H22) spray flame in a laminar counterflow field, with a burner setup and flame pictures shown in Figure 48 [603]. A vibratory orifice atomizer (VOA), installed at the top of the upper (fuel) burner, provided quasi-monodispersed fuel droplets with variable mean droplet sizes. The fuel spray was premixed by air at $\phi = 0.106$ and issued from the upper nozzle against stretch-stabilized lean methane-air premixed flames, mimicking the conditions in practical combustors where premixed fuel sprays approach the high temperature zone. A strong effect of initial droplet size on soot formation was observed. As the initial Sauter mean diameter (SMD) of the fuel droplets increased, an initially blue reaction zone transitioned towards yellow flames with increasingly higher luminosity, indicating an enhancement of both the peak soot concentrations and the sooting zone areas (further confirmed by corresponding LII measurements). In addition, an interesting local inhibition of soot formation was observed for larger SMD conditions, resulting in an annular shape for the spatial distribution of soot, with particle sizes being larger in the radially outer region of the soot zone. This local inhibition can be attributed to the evaporative cooling effects caused by group combustion of the unburned, initially large fuel droplets.
Figure 48. Experimental setup for the investigation of counterflow flames of fuel sprays employed by Hayashi et al [603]. Shown are the counterflow burner with two opposing converging nozzles, the vibratory orifice atomizer attached to the upper burner for the generation of liquid sprays with controllable droplet sizes, and a schematic of the spray CDF. Reprinted from Hayashi et al [603] with permission of Elsevier.

Droplet size distribution can also significantly affect soot formation. Experimental results showed that both the soot concentration and the sooting zone area were reduced in spray counterflow flames for poly-dispersed droplets, as compared to quasi-mono dispersed droplets with the same SMD [612]. In poly-dispersed cases there were a larger number of small fuel droplets, more likely to be completely evaporated before reaching the high temperature flame zone. As a result, the premixed-like combustion mode was promoted, which is helpful for soot reduction.
5.6.2 Effects of electric fields on soot formation in CDFs

Hydrocarbon flames are abundant with electrically charged species [830] such as the \( \text{H}_3\text{O}^+ \), \( \text{CHO}^+ \), \( \text{CH}_3^+ \) and \( \text{C}_3\text{H}_3^+ \) ions [831, 832], ions of PAHs [833], and carbon clusters in the form of \( \text{C}_{2n} \) with \( 30 \leq 2n \leq 210 \) (both positive and negative) [834, 835], in addition to charged soot particles [133, 836, 837]. It is therefore reasonable to expect the flame to be significantly affected when exposed to external electric fields. Indeed, it has been shown that electric field-assisted combustion can change flame shapes [838, 839] and temperatures [840], improve flame instability [841-843], enhance flame propagation speeds [844-846], and affect soot emissions [527, 566, 847-850].

Whether soot formation in a certain CDF can be inhibited depends on the intensity, polarity (for DC), and frequency (for AC) of the applied electric fields [527]. In sooting ethylene CDFs, where the fuel nozzle is connected to the high-voltage terminal and the oxidizer nozzle is grounded, a small positive voltage did not notably affect the flame position or soot formation, but larger voltages resulted in flame instability. However, applying negative voltage \(< -2\text{kV}\) reduced soot/PAH concentrations significantly in both SF and SFO flame configurations. The measured spatial distributions of OH / PAH LIF, and soot LII signals, are shown in Figure 49 for both the SF (a) / SFO (b) baseline flames and those with a negative voltage of \(-2.0\ \text{kV}\) applied (c, d). The locations of the OH LIF zones were seen to be dragged towards the fuel side by the electric field, consistent with the direction of ionic wind [851, 852], since there are more positive ions than negative. Similarly, the stagnation plane was also forced closer to the fuel side. Since soot is always formed on the fuel side of the flame, these movements of the flame front and stagnation plane are expected to result in a narrower fuel pyrolysis zone and less residence time for PAH/soot growth. Furthermore, heat loss toward the burner nozzle may also be enhanced, leading to lower flame temperature and reduced soot formation rates. Note that these explanations are based solely on the effects of ionic wind. Although reasonable, they do not explain the flame instability when positive potential is applied, or the fact that
in certain flames [850] luminosity was always seen to shift away from the fuel nozzle, independent of the polarity of the DC potential.

![Diagram](image)

**Figure 49.** Effects of DC electric field on measured spatial distributions of OH / PAH LIF and soot LII signals. Reprinted from Park et al. [527] with permission of Taylor & Francis.

Kono and coworkers [850] applied AC potentials in sooting C₂H₂ CDFs. At an AC frequency of 327 kHz, the authors observed that SVF profiles shifted away from the fuel nozzle and peak SVF increased when the peak-to-peak voltage was 4.8 kV, while the opposite was true when it was 1.6 kV. These complex behaviors were thought to result from forces other than the ionic wind effects. Clearly, more mechanistic investigations are required to explain these experimental observations.

### 5.6.3 Effects of soot radiation on flame characteristics

CDF is a classic tool for investigating critical strain rate in diffusion flame extinction [109, 853], which are routinely used as an important global flame property for the validation of detailed kinetic
mechanisms (see e.g., Refs. [587, 854-856]). CDF can be extinguished at both high and low strain rates [857-860]. Available time for mixing is short at sufficiently high strain rates, and the flame is extinguished due to insufficient fuel/oxidizer mixing. At very low strain rates, flame intensity is low and relative heat loss due to radiation becomes critically important, which can notably decrease flame temperature (increase reaction time) and finally, extinguish the flame.

Heat loss due to radiation from soot particles can be significant, especially when considering the multi-dimensional effect in practical flames where soot produced in fuel-rich regions can be transported (by convection and/or turbulence) to the high-temperature reaction zone to exert influence on flammability. To analyze the effects of radiation heat loss from this non-local soot on the flame extinction limit, an analytical study, using large activation energy asymptotic theory, was performed for C$_2$H$_4$ CDF, with pre-loaded soot particles in the oxidizer stream [604]. The results, shown in Figure 50, demonstrate that the presence of soot had a strong effect in reducing the size of the flammable domain. Particularly, with an initial soot mass fraction of 5%, the lower strain rate radiation-extinction limit would be increased by more than one magnitude, and the upper mixing-limited extinction strain rate could be decreased by around 30%.

In vigorously burning CDFs, the presence of flame-generated soot particles also tends to reduce soot formation as a result of radiative cooling of the flame [583, 613, 627, 861]. The extent of such reduction is higher in flames with a lower strain rate [627]. Nevertheless, in typical SF flames, the effect of soot radiation was found to be weaker than that of gas radiation [583]. This is because in SF flames, SVF peaks at the stagnation plane, which is spatially separated from the high-temperature reaction zone and thus has a relatively low temperature of around 1000 K. On the other hand, the most effective gaseous radiators such as CO$_2$ and H$_2$O peak near the reaction zone with much higher local temperature. For the same reason, soot radiation in SF flames may be less important than in coflow flames, where SVF peaks in regions of relatively higher temperatures.
Figure 50. Effects of soot loading on extinction strain rate in counterflow diffusion flames, numerical data taken from Narayanan et al. [604]. For non-adiabatic flame, extinction can be seen to occur at both high and low strain rates due to an excessively small Damköhler number; and the presence of soot particles noticeably affects the lower critical strain rate for extinction.

5.6.4 Interactions between soot and nitric oxides

The formation of nitrogen oxides (NOx) in hydrocarbon-air flames is known to be temperature sensitive and can therefore be influenced by radiative heat loss from gaseous species and soot particles, if present [862, 863]. In addition, oxidation of soot particles and precursors may compete with thermal NOx formation for O and OH radicals, contributing to the reduction of NOx emission [539]. On the other hand, nitric dioxide (NO2), as a form of NOx, is a strong soot oxidizing agent—even more effective than O2 [327, 330]. In this regard, soot and NOx—being the two most worrying pollutants from hydrocarbon fuel combustion—can interact in their formation processes.

A dedicated numerical analysis was performed to investigate soot and NO interactions in ethylene CDFs of the SF type [864]. The model included a detailed reaction mechanism with NOx chemistry, a two-equation semi-empirical C2H2-based soot sub-model [376], and radiation heat loss from major gaseous species and soot particles. Three simulation runs were conducted to investigate the detailed mechanisms that determine how the presence of soot affects NOx formation. In the first run (SIM1), both NOx and soot models were included, and in the third run (SIM3) only the NOx model was
The second run (SIM2) was computed with a fixed temperature, obtained from SIM1, while all rates related to soot were set to zero. As a result, the difference of the predicted NOx concentrations between SIM2 and SIM3 was caused by purely thermal effects from soot radiation. The differences between SIM1 and SIM2 were due to the chemical effect of soot formation, as the production of soot is expected to alter the chemical structures of the flames by—for example—consuming gaseous soot precursors. Both thermal and chemical effects of soot on NO formation were observed, as can be seen in Figure 51. In terms of the chemical effect, analysis of the detailed reaction pathway suggests that the most important competing species between soot and NOx chemistry is C2H2.

On the other hand, comparison between two numerical runs with and without the NOx sub-chemistry shows that the formation of NOx has a negligible effect on soot formation/oxidation in the flames investigated. However, it should be noted that the authors [864] only considered the effect of NO, and further investigations taking NO2 formation into account may be needed. In addition, studies on more flame configurations, like SFO CDFs, are necessary to confirm whether the insignificance of NOx on soot formation can be generalized. This is especially relevant considering that in SF CDF, the peak locations of NOx and soot are, in fact, spatially separated.

![Figure 51](image.png)

**Figure 51.** Effects of soot formation on NO formation in CDFs. Data taken from Guo *et al.* [864].
6. Summary, challenges, and prospects

In view of the growing global energy demand, conventional hydrocarbon fuel is expected to continue as the dominant energy source in the foreseeable future. However, the consensus is that hazardous emissions from the combustion of hydrocarbon fuels must be mitigated for a sustainable future. As a ubiquitous combustion phenomenon, soot formation has attracted continuous research attention, not only because of its associated negative environmental impact, but for the rich physicochemical processes involved which are still not fully understood. Nevertheless, the outcome from decades of fundamental research—especially those performed in well-controlled laboratory-scale flames—confirms that it is possible to describe soot formation from first principles.

Although the primary objective of this review is to provide a comprehensive summary of soot research performed with counterflow flames, many relevant aspects involve fundamental soot mechanisms that are universally applicable and cannot be isolated from other research. Therefore, the concluding remarks that follow list several general challenges and opportunities for soot research, before discussion of those more specific to counterflow flames.

As discussed extensively in the text, the conceptual stages of soot formation include 1) gas phase molecular growth towards large PAHs; 2) soot nucleation out of the gas phase; 3) surface growth and particle-particle coagulations; and 4) soot oxidation and fragmentation. Despite research progress made thus far, there are still unresolved aspects in each of these processes that continue to challenge the soot research community. The most prominent challenges (in our opinion) are as follows:

- Molecular growth from aliphatic fuels involves the formation of the first aromatic ring, in most cases, benzene. Although there are various reaction pathways for benzene formation, and the relative importance of each pathway may depend on fuel molecular structure, the number of pathways is finite, and fairly detailed knowledge is available about many of them [172]. However, the story is far different for molecular growth beyond benzene. As discussed in section 2.1.2, new pathways for PAH growth are being identified (and proposed) continuously.
But experimental validation of these pathways becomes challenging since a) the concentration of large PAHs are typically too low for accurate measurement; and b) experimental techniques to resolve the many isomers of large PAHs are limited. In fact, in view of the formidable number of isomers for molecules with increasingly larger sizes, the question remains whether all the relevant pathways [865] can ever be captured. A relevant question is to what details would we need for practical purposes.

- The transition from gas-phase molecular soot precursors to condensed-phase particles has remained a mystery for many decades. The assumption of physical dimerization of PAHs has been implemented in many soot models, successfully explaining many experimental observations. However, theoretical analysis determined that at flame temperatures, PAHs with enough abundances are too volatile to be physically linked. As such, it is believed that covalent interactions are needed to bind the PAHs to form molecular clusters (incipient particles). While previous mechanisms for covalent cluster formation required repeated high energy barrier activation of stable PAHs (and thus not perfectly compatible with the rapid nucleation of soot, even in low temperature radical-starved regions), the recently proposed clustering of hydrocarbon by radical chain reaction (CHRCR) mechanism [138] highlights the role of radical chain propagation through resonantly stabilized radicals. This new CHRCR mechanism overcomes the issue of low molecular growth rates associated with existing chemical nucleation mechanisms and also allows σ-dimerization from large RSRs [138, 273]. Although promising, one challenge is that detailed reaction rates must be established for the proposed reactions so that an extensive kinetic mechanism can be assembled to test whether CHRCR can explain experimental features on soot formation. The target for testing may include, for example, the bimodal size distribution in the post-flame zone of a premixed stagnation flame, the non-monotonic variations of soot formation with temperature in a shock tube and premixed flames, the synergistic effects between fuels on soot formation in diffusion flames and many
more. Furthermore, the role of charged ions in soot nucleation must be explored further; it is possible that different mechanisms can simultaneously contribute to soot nucleation.

- Reactions that occur on the surface of soot particles account for most of soot mass growth, so a deeper understanding of these surface reactions is critical in order to predict the amount of soot formed in flames and practical combustion devices. In most current soot models, surface reaction is dominated by the surface-HACA mechanism, in which C$_2$H$_2$ is the most important mass growth agent. Rates for heterogeneous hydrogen abstraction and C$_2$H$_2$ addition reactions are obtained by assuming a kinetic analogy between soot and gas phase aromatic species. Currently, however, it is unclear how and to what extent surface morphology, adsorption/desorption and other features specific to heterogeneous reactions will affect these surface reactions [865]. The effects of surface properties (e.g., graphitization) on soot growth are yet to be studied quantitatively. To date, there is no fundamental models that describe how the surface properties change as the soot particle evolves in the flame. Such lack of detailed knowledge about surface reactions is reflected in the wide variability of the parameter $\alpha$ (representing the fraction of surface site that is active for surface reactions) in different soot models. It is a common practice to adjust $\alpha$ to match experimental soot concentrations. In addition to C$_2$H$_2$, the recent CHRCR mechanism [138] also suggests that surface growth can proceed with radical-chain reactions through RSRs. The possibility that aliphatic chains can be present at the surface of nascent particles adds more complexity to a complete description of surface reactions. It is obvious that more fundamental research is required to gain further insights into the heterogeneous reactions on soot surfaces.

- Soot oxidation is essential in determining ultimate soot production, and in the regeneration of particulate filters, which are now routinely used to control soot emission from IC engine-powered ground vehicles. The oxidation of soot involves complex heterogeneous surface reactions; therefore, many of the challenges associated with soot surface growth discussed
above also apply to soot oxidation. In addition, experimental evidence of the particle internal burning mode [348] may also suggest the importance of surface diffusion of oxidizing agents. However, many current models for soot oxidation are still largely empirical in that they typically take bulk carbon as the sole solid reactant, neglecting any steric effects and the many different kinetic features of soot formed in different environments. Intermediate products of soot oxidation also require further research attention as it was shown in SFO flames, the oxidation of soot may produce a second peak for PAH LIF signals, indicating the complexity of soot oxidation products. It is clear that future research is needed to uncover more detailed and stERICally-resolved information on how oxidizing species such as O, OH, O2 and NO2 interact with soot particles for the removal of soot mass.

In addition to these challenges, soot formation is further complicated because of its dependence on many combustion parameters. The present review clearly demonstrates that research conducted in counterflow flames has notably contributed to our understanding of the effects of fuel molecular structure, fuel/oxidizer dilution, pressure, temperature, residence time, flame structure, and unsteadiness on soot formation. Further investigations of the underlying mechanisms of these phenomenological effects in turn provide hints through which various aspects of soot formation fundamentals can be uncovered. This review also identifies several challenges that must be addressed for more complete soot characterization in counterflow flames and for further contributions to a first principle understanding of soot. Following are some pertinent issues, along with prospects for future research:

- Characterization of soot in counterflow flames is typically limited to soot volume fraction, primary particle size, and total number densities, while more detailed information such as particle size distribution (PSD) is rarely reported. This is because conventional sampling methods are too intrusive for counterflow flames, which have a very narrow sooting zone (~1 mm). However, PSD can provide valuable information about nearly all aspects of soot
information and it is essential if detailed insight is to be derived from counterflow flame experimental data. Future effort is needed to develop reliable, *in-situ* methods (e.g., time-resolved LII) to measure PSD in CDFs. Challenges also exist for the thermophoretic extraction of soot particles in counterflow flames, because the TEM probe used in traditional fast-insertion methods may vibrate too much to allow for soot extraction at a specific location. The SiC wire method developed by Figura and Gomez [508] may partly solve this problem.

- The narrow sooting zone in counterflow flames also indicates that complete soot evolution from gas-phase soot precursors (via nascent particles to mature soot particles) occurs in a somewhat confined spatial region, necessitating high spatial resolution experimental techniques. Conventional optical methods for quantitative soot measurements, such as light extinction and scattering, are dependent on the complex refractive index of soot, which is known to vary with particle size and maturity. However, a quantitative relationship is yet to be determined. Additional methods may be necessary (i.e., small angle X-ray or neutron scattering [866, 867]) to confirm the extent to which the varying optical properties may affect the quantitative soot measurements in counterflow flames.

- There exist a number of counterflow flame-based studies regarding the synergistic effects of fuel mixtures on PAH and soot formation, demonstrating that certain binary fuel mixtures can have higher sooting tendency than either fuel component (e.g. [86, 528, 586]). These works highlight the importance of chemical interactions between different fuels and so have practical application for fuel formulation. While future opportunities exist for the identification and kinetic explanation of additional fuel pairs exhibiting synergistic effects, it is interesting to search fuel mixtures with lower sooting tendencies than either component, which exhibit a negatively synergistic effect.

- To be more relevant to practical combustion systems like diesel engines and jet engines, counterflow flame experiments must be extended to operate with liquid fuels at high-pressure
conditions. Although the counterflow configuration is advantageous for controllable and stable flame operation at elevated pressures (i.e., not sensitive to buoyancy-driven instability, no heat loss through burner rim) [788], it is important to design boundary conditions carefully, so that pressure effects on soot formation can be isolated [537, 618, 788]. In terms of fuel types, studies on oxygenated fuels with aromatic rings are scarce, but they deserve attention in future research because of their practical uses and potentially different soot growth pathways, as compared to other aliphatic and aromatic fuels.

– Counterflow flames are particularly suitable for studying the effects of fuel-borne catalysts on soot formation and oxidation (FBCs, like those metal-containing compounds reviewed here). This is because the formation and oxidation of soot can be separated by utilizing either SF or SFO flames. The challenge is to develop effective measurement techniques that distinguish soot and catalytic particles so that concentrations of each can be reliably determined. Possibilities include methods based on spectral emissions [868, 869], but it should be considered that these methods need high spatial resolution for applications in counterflow flames.

– Time-resolved experimental data on PAH, and soot in CDFs subject to oscillating strain rates, are scarce [585, 801]. Such data can provide not only information about the unsteady flamelet model, but it can validate kinetic models for PAH and soot formation. Peaks, troughs, and mean values of PAH/soot concentrations in oscillating flames, normalized by their steady-state values, provide valuable information regarding the kinetics of their formation.

Acknowledgements

The preparation of this manuscript was supported by the National Natural Science Foundation of China (51606136) and the Clean Combustion Research Center of KAUST. The authors are grateful to Drs. S. Mani Sarathy, Hong G. Im and William L. Roberts of KAUST, Dr. Abhijeet Raj of Khalifa University and Dr. Sungwoo Park of the Korea Aerospace University for collaboration on relevant
projects. YW also acknowledged stimulating and informative discussions with Dr. Alessandro Gomez of Yale University, Dr. Dongping Chen of Beijing Institute of Technology and Dr. Xunchen Liu of Shanghai Jiaotong University. Technical assistances from Lei Xu, Wei Dai, Mengxiang Zhou and Wei Wang of Wuhan University of Technology in organizing some of the cited references are also acknowledged.

References:

[90] Sirignano M, Salamanca M, D'Anna A. The role of dimethyl ether as substituent to ethylene on particulate formation in premixed and counter-flow diffusion flames. Fuel 2014;126:256-62.


[520] Lehtipalo K. How the development of condensed particle counters is reforming our view on atmospheric nucleation. Helsinki, Finland: University of Helsinki; 2011.


[781] Mandatori PM, Gülder ÖL. Soot formation in laminar ethane diffusion flames at pressures from 0.2 to 3.3 MPa. Proc Combust Inst 2011;33:577-84.


